Elasticity, viscoelasticity, and glass transition of model systems by computer simulation

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Spécialité: Physique

ELASTICITY, VISCOELASTICITY, AND GLASS TRANSITION OF MODEL SYSTEMS BY COMPUTER SIMULATION

par

Da LI

Soutenue le 24 novembre 2016 devant le jury composé de:

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Abstract

Elasticity, viscoelasticity and glass transition of glass-forming liquids and model solids are investigated by computer simulations at equilibrium using the stress fluctuation formalism. The model systems studied include Lennard-Jones systems at two and three dimensions, a binary mixture of Lennard-Jones (LJ) particles constituting the two-dimensional version of the well known glass-former Kob-Andersen model (KA2d model), an AB13 binary mixture of repulsive particles and a ternary mixture of hard spheres. Our studies aim mainly at the effect of the temperature $T$ on the mechanical properties of the systems under investigation, with special focus on phase transitions such as the crystallisation, or the glass transition. Our results are presented in two parts, namely static properties and time-dependent properties. The static properties we studied are essentially the shear modulus $G$ and the bulk modulus $K$. Using molecular dynamics simulations provided by the LAMMPS code, we characterized $G$ and $K$ of the glass-former KA2d model, and compared the results with a one-component LJ2d system. While the latter shows a significant jump in $G$ at the freezing transition, the former displays a continuous cusp-type behaviour at the glass transition $T_g$, following the law $G(T) \propto (1 - T/T_g)^a$ with $a \approx 0.6$, confirming qualitatively previous simulation results carried out on KA3d system in 2013, but in contradiction to predictions based on mode-coupling theory. We also compared the elastic constants of an LJ fcc crystal with those of a more complex solid, the AB13 structure. This study shows that while the non-affine contribution vanishes at zero temperature for the fcc structure, it is not the case for the AB13 structure. This finding points out that we cannot use systematically the approximation of affine elasticity at $T = 0$ for crystals of more than one component. In the part devoted to time-dependent properties, we investigated several aspects and functions. First, we explored the sampling-time dependence of our static
results, and show that this influences strongly the stress fluctuations. Second, equilibrium dynamics are studied via the shear-stress autocorrelation and the shear-stress mean square displacement, and the relation between them. The main focus, however, is the shear-stress relaxation modulus $G(t)$ and the associated dynamic moduli $G'(\omega)$ and $G''(\omega)$, with $\omega$ the frequency. These functions characterize the viscoelasticity of our systems. Using a recently proposed formula (Wittmer et al., Phys. Rev. E 93 2016), we are able to determine these functions quite accurately and study their behaviour in liquid, crystal and glassy states.
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Chapter 1

Introduction

This thesis investigates mechanical properties such as elasticity, viscoelasticity, and glass transition of model systems by means of molecular simulations.

Elasticity is one of the fundamental properties of materials. It shows the deformation capability of a substance under the external force field. In macroscopic scale and in the linear response regime, Hooke's law is used to characterize the elasticity of materials, by introducing the elastic constants. With the development of simulation techniques, we are able to link microscopic aspects of materials with these macroscopic constants. The fluctuation formalism method is one of them, in which the elastic constants are related to the fluctuations of the stress tensor components. This approach was first introduced by Squire et al. [1] to calculate the isothermal elastic constants for argon crystal at finite temperatures by Monte Carlo (MC) simulations. The formalism was also used in Molecular Dynamics (MD) simulations by Barrat et al. [2] to calculate elastic response of a binary alloy near the glass transition. The remarkable advantage of this approach is that the unstrained systems are used, thus the elastic constants can be obtained via equilibrium molecular simulations. In reference [1], the constants are expressed by the sum of three contributions: the kinetic term, the Born term and the stress-fluctuation term. The kinetic term is the ideal gas term. It is zero for $T = 0$. The authors suggested that the fluctuation term arises only with non zero temperatures. Thus at zero temperature, only the Born term would subsist, leading to the well known formulas by Born and Huang [3] for the elasticity of solids at $T = 0$. The issue was later discussed by Lutsko [4], who pointed out that even at $T = 0$, the fluctuation term can be non zero if the solid is complex, for example
having more than one atom per primitive unit cell. The Born term accounts for the average stress resulting from an affine uniform microscopic strain field of the system in response to a macroscopic strain. The fluctuation term, at $T = 0$, corresponds hence to non-affine displacements of the particles in order to relax the internal stress. At $T > 0$, it represents indeed the thermal fluctuations of the stress tensor elements. This prediction was verified for model glassy systems at $T = 0$, where the fluctuation term was found as important at half the magnitude of the Born term [5]. Another interest of the method is that calculations can be done for $P \neq 0$ [6], making it an attractive tool to study the elasticity of materials for $T \neq 0$, and $P \neq 0$. In crystals, there are usually more than two independent elastic constants [7]. For isotropic bodies, there are only two of them. The convenient set can be the so-called elastic moduli [8], consisting of the bulk modulus $K$ and the shear modulus $G$. Their macroscopic definitions are well known: $K = -(1/V)(\partial P/\partial V)_T$ and $G = \delta \tau / \delta \gamma$, where $\delta \tau$ is the average shear stress and $\delta \gamma$ a small shear strain. Their “microscopic” expressions are well defined too within the stress fluctuation formalism and shall be explained in detail in the thesis.

Although this formalism was initially proposed for solids, the formulas for computing the shear modulus $G$ and the bulk modulus $K$ are found to be valid for the liquid state, where one obtains consistently $G = 0$ and $K$ being the same expression as the one obtained by Rowlinson in the context of an isotropic liquid state [9]. This observation is important as one studies the liquid-solid phase transition, as we do in this thesis. We are thus able to use the stress fluctuation formalism on both phases and through the transition. Beyond characterizing $G$ and $K$ through a liquid-crystal transition, we are also interested in low temperature crystals. We shall verify indeed that for simple crystals, the non-affine contribution to the elasticity is zero, whereas for more complex crystals, such as the AB13 system [10], this contribution is on the other hand non-zero, as suggested by Lutsko.

All materials display some viscoelastic response. The mathematical formulation of viscoelasticity theory makes it possible to predict the material response to any load history. Viscoelasticity is beneficial in a number of fields of materials science, metallurgy, and solid state physics as it is causally connected to various microphysical processes and can be used as an experimental probe of those processes [11].
links between viscoelasticity and microstructure are exploited in the use of viscoelastic tests as an inspection tool as well as in the design of materials [11]. In complex fluids, viscoelasticity is also an important property [12, 13]. It can reveal different time scales involved in the stress relaxation process. It is thus a precious probe of dynamics of complex fluids. The shear relaxation modulus $G(t)$ gives the measure of the viscoelasticity of a system. This function can be measured in rheological experiments. More usually, the dynamic moduli $G'(\omega)$ and $G''(\omega)$ (with $\omega$ the frequency) are measured, thanks to oscillatory stresses. As $G'$ and $G''$ are related to $G(t)$ in a simple way via sine and cosine Fourier transformations, knowledge of one set leads to the other set.

As we know, computer simulations can bring a lot for the understanding of materials properties. It is important that we can compute these functions by using statistical physics tools. In liquids, it is well known [14] that $G(t)$ is given by the shear stress autocorrelation function (SACF), which can be computed by equilibrium molecular dynamics simulations. For example, recently Heyes et al. [15] investigated the elastic moduli, viscoelasticity properties of the Lennard-Jones fluid along the fluid-solid coexistence line by equilibrium Molecular Dynamics simulation. For solids, however, generally one cannot obtain $G(t)$ directly from the SACF [16]. Recently, a formula [17] has been proposed and tested for various model systems. It has been proven to be robust and general for systems as different as permanent elastic bodies [17], or self-assembled transient networks [18]. In this thesis, we shall apply this formula to our model systems. Our aim is to compare the dynamic moduli in various states: liquids, crystals and glasses, in order to reveal some general trends, related to different states.

The third topic of the thesis is the glass transition. Generally speaking, the glass transition is a transition by which a liquid changes into an amorphous state without crystallization. The most ordinary way of glass transition is quenching, namely, supercooling a liquid. By this method, lots of glassy substances can be obtained. The remarkable phenomenon upon supercooling is that the viscosity increases dramatically from the liquid side to glass. The glass is similar to the relevant liquid in structure. In terms of mechanical properties, it is solid-like. The glass transition has attracted much attention. Many models and theoretical approaches are proposed to explain it, such as Adam-Gibbs model, Kob-Andersen model, random first order transition theory, and
mode coupling theory. In opposition to the freezing transition, the glass transition is continuous in specific volume change and the transition temperature $T_g$ is cooling-rate dependent. The Kob-Andersen (KA) model has been investigated extensively since it was first proposed to test mode-coupling theory [19]. The KA model is a binary mixture of Lennard-Jones (LJ) particles. Another popular glass-former is the polydisperse LJ systems [5]. With these LJ glasses, different aspects are studied, such as static yield stress [20], aging effects [21], fluctuation dissipation ratio [22], single particle jumps [23], local elasticity map and plasticity [24]. The examined assumption impact factors of glass transition contain static length scale [25], point-to-set length scales [26], heterogeneity [27]. As we can see, the elasticity is not enough investigated and we focus our investigations on this topic. As is well known, the shear modulus is zero in liquids and non zero in solids (crystal or glass). This quantity can thus be a signature of the glass transition, as an order parameter. This is for example the criterion used in [28]. In this case, one can question about the nature of the change in $G$ at $T_g$, whether it is a continuous change, or a jump. Since there is still debate on this question, we carry out investigations on KA2d model system with focus on this topic.

In order to examine the above mentioned properties using the stress fluctuation formalism, we choose a few simple model systems in this thesis. The glass transition is especially studied with the KA2d system. The companion system, the monodisperse LJ2d system, which crystallizes, is also studied for the sake of comparison. Two crystals are investigated for their elasticity and viscoelasticity, the first one is the monodisperse LJ3d system which forms a fcc crystal. The second is the AB13 system which forms a cubic complex superlattice. These studies are done by using Molecular dynamics simulations provided by the LAMMPS code [29]. To a minor extent, monodisperse and polydisperse hard sphere fcc crystals are also studied by constant pressure Monte Carlo simulations [30], in order to test the computation of the bulk modulus by volume fluctuations.

The thesis is organized as follows: in chapter 2, we present the methodology used in this thesis. After an overview of classical simulation methods, Monte Carlo and Molecular Dynamics, we discuss the different statistical ensembles that are relevant
to our studies. They are microcanonical (NVE), canonical (NVT) and isothermal and
isobaric (NPT) ensembles. A large portion of this chapter is devoted to the presentation
of the stress fluctuation formalism for the calculation of the elastic constants. Chapter 3
contains our results on static properties of the model systems we studied. Three topics
have been considered. The first is the calculation of bulk moduli of monodisperse and
polydisperse hard sphere crystals by volume fluctuations, the second is a study of a
glass former in two dimensions: 80-20 Kob-Andersen model, and the last topic concerns
shear stress fluctuations in simple and complex crystals. Chapter 4 investigates time-
dependent properties of our systems. Several questions have been examined. The
first is the sampling-time dependence of the stress fluctuations, then the shear-stress
autocorrelation function and the shear-stress mean-square displacement are considered.
The last part concerns the viscoelastic properties of our model systems, in liquid, glass
and crystal states. Conclusions are gathered in chapter 5.
Chapter 2

Methodology

In this chapter we shall introduce the theoretical foundation of our work. First we shall describe briefly the simulation methods, Monte Carlo (MC) and Molecular dynamics (MD) simulations. A brief discussion on the relevant statistical ensembles is given. Then we shall present the method of computing the elastic properties of a model material using the stress fluctuation formalism.

2.1 Monte Carlo methods

Monte Carlo methods used in numerical analysis are mainly aimed at calculating multivariable integrals. In molecular simulations, it is used to simulate statistical systems and produce a representative ensemble of configurations to reach thermodynamical quantities without the necessity (or the possibility) of computing them analytically [31].

2.1.1 Monte Carlo integration

Monte Carlo technique can be a method of integration. Take a simple integral for example

\[ F = \int_{x_1}^{x_2} dx f(x). \]  

(2.1)

This integral can be calculated approximately by

\[ F' = \frac{x_1 - x_2}{N} \sum_{i=1}^{N} f(x_i) \]  

(2.2)
where the points \( x_i \) are evenly distributed in the range \((x_1, x_2)\). As long as we select enough points, that is, the number \( N \) is big enough, the \( F' \) will be sufficiently close to the exact \( F \). However, the points can be also selected randomly and in this case the integral can be written as follows

\[
F = \int_{x_1}^{x_2} dx \frac{f(x)}{\rho(x)} \rho(x)
\]

with \( \rho(x) \) an arbitrary probability density function \((\rho(x) \neq 0)\). Consider conducting lots of trials \( tr \), and every time a random number \( \xi_{tr} \) is chosen from the distribution \( \rho(x) \) in the interval \((x_1, x_2)\). We obtain that

\[
F = \langle f(\xi_{tr}) \rangle_{trials}
\]

where the brackets show an average over all trials. The probability density function \( \rho(x) \) could be as simple as a uniform distribution (sample mean integration),

\[
\rho(x) = \frac{1}{x_2 - x_1}, \quad x_1 \leq x \leq x_2,
\]

following this, the integral can be approximated by

\[
F \approx \frac{x_2 - x_1}{tr_{max}} \sum_{tr=1}^{tr_{max}} f(\xi_{tr}),
\]

where \( tr_{max} \) is the total trial number. We notice that for a one-dimensional integration, this method is not as efficient as many others (Simpson, Gauss, etc). But its use is more indicated for multi-dimensional integrations.

### 2.1.2 Importance sampling

In Monte Carlo algorithms, lots of trials may give a very small contribution to the Monte Carlo average if we use an uniform sample mean integration method (above). Importance sampling techniques can solve this problem, by choosing random numbers from a non-uniform distribution [30]. They concentrate the samplings in the regions of interest which make important contributions to the integral and hence enhance the overall efficiency. This technique is essential to molecular simulations. For example, in the canonical ensemble \((NVT)\), if we want to calculate the average of a physical quantity \( A \), we write

\[
\langle A \rangle_{NVT} = \int d\Gamma \rho_{NVT}(\Gamma) A(\Gamma),
\]

\[7\]
where $\Gamma$ is a point of the phase space, and $\rho_{NVT}(\Gamma)$ is the canonical distribution. Comparing with eq.(2.1), we have $f = \rho_{NVT}\mathcal{A}$. In general, the integrand $f$ will be important where $\rho_{NVT}$ is important, and neglectable where $\rho_{NVT}$ is neglectable. By repeating the previous MC integration method we obtain the integral

$$
\langle \mathcal{A} \rangle_{NVT} = \langle \mathcal{A} \rho_{NVT}/\rho \rangle_{\text{trials}}. \tag{2.8}
$$

if we choose $\rho = \rho_{NVT}$, we perform an efficient estimate of the integral. Then

$$
\langle \mathcal{A} \rangle_{NVT} = \langle \mathcal{A} \rangle_{\text{trials}} \tag{2.9}
$$

In order to generate a series of random states at the end of the simulations where every state has appeared with the appropriate probability, a Markov chain of states of the system is set up. This is done using the Metropolis algorithm below.

**2.1.3 The Metropolis method**

The Metropolis algorithm is used in molecular simulations. It generates a sequence of states according to the equilibrium probability distribution of a given system. The following gives a presentation according to Chandler’s book [32]. We only consider the change of particle’s positions. Suppose a state $m$ of the system, and $\rho_m$ the equilibrium probability, consistent with the Boltzmann distribution $\rho_m \propto \exp (-\beta \mathcal{V}_m)$, ($\beta = 1/k_B T$, and $\mathcal{V}_m$ is the potential energy of the system at state $m$). If $T_{mn}$ is the transition probability (per unit time) that the system transit from state $m$ to $n$, then the kinetic equation for $\rho_m$ writes

$$
\dot{\rho}_m = \sum_n (\rho_n T_{nm} - \rho_m T_{mn}). \tag{2.10}
$$

At equilibrium, we have $\dot{\rho}_m = 0$ (the equilibrium state is a stationary solution of the kinetic equation). From eq.(2.10), we see that this is the case when

$$
\rho_n T_{nm} = \rho_m T_{mn}. \tag{2.11}
$$

Eq.(2.11) is referred to as the detailed balance condition. More precisely, we can write

$$
\frac{T_{mn}}{T_{nm}} = \frac{\rho_n}{\rho_m} = \exp (-\beta \Delta \mathcal{V}_{mn}) \tag{2.12}
$$
where $\Delta \mathcal{V}_{mn} = \mathcal{V}_n - \mathcal{V}_m$. In practice, the Metropolis Monte Carlo (MMC) scheme proposes

$$T_{mn} = \begin{cases} 
\alpha_{nm} & \text{if } \rho_n \geq \rho_m, \\
\alpha_{nm} \frac{\rho_m}{\rho_n} & \text{if } \rho_n < \rho_m.
\end{cases} \quad (2.13)$$

where $\alpha_{mn} = \alpha_{nm}$ being the elements of a symmetric stochastic matrix. Here is an MC step: suppose the system at state $m$, a trial move is made, by displacing one atom randomly, tempting to bring the system to state $n$. If $\Delta \mathcal{V}_{mn} \leq 0$, then the move is accepted. If $\Delta \mathcal{V}_{mn} > 0$, then $\exp (-\beta \Delta \mathcal{V}_{mn})$ is compared to a random number $\xi$ picked up uniformly in $[0, 1]$. The move is only accepted when $\exp(-\beta \Delta \mathcal{V}_{mn}) > \xi$. In other words, in the Metropolis algorithm, the acceptance probability for a move from state $m$ to state $n$ is

$$P(m \to n) = \min \left(1, \exp \left(-\beta (\mathcal{V}_n - \mathcal{V}_m)\right)\right) \quad (2.14)$$

It is important that the Markov chain generated in such a fashion is ergodic. That means from any state, any other state should be reached after a finite number of trial moves. An MC simulation contains a very large number of MC steps, typically more than a million.

### 2.1.4 MC simulation of hard spheres

The hard sphere (HS) model is a simple model to represent dense fluids. The HS potential is

$$\beta u(r) = \begin{cases} 
0 & \text{if } r \geq \sigma \\
\infty & \text{if } r < \sigma
\end{cases} \quad (2.15)$$

where $\sigma$ is the HS diameter. This singular potential leads to a very simple Metropolis MC algorithm. Consider the system be in state $m$. The state $n$ consists of displacing one particle $i$ a small distance: $r_i^{(m)} \to r_i^{(n)}$. Then, substituting eq.(2.15) in eq.(2.14), the acceptance probability for a move from state $m$ to state $n$ is

$$P(m \to n) = \begin{cases} 
1 & \text{if } \left|r_i^{(n)} - r_j^{(m)}\right| \geq \sigma \text{ for all } j \neq i \\
0 & \text{if } \left|r_i^{(n)} - r_j^{(m)}\right| < \sigma \text{ for one } j \neq i,
\end{cases} \quad (2.16)$$

i.e. it is an overlap test. The scheme is easily generalized to HS mixtures, where each species has a diameter $\sigma_\mu$. If the particle $i$ belongs to species $\mu$, its position is $r_{i\mu}$. 

9
The MC acceptance probability is then

$$P(m \rightarrow n) \begin{cases} 1 & \text{if } |\vec{r}_i^{(n)} - \vec{r}_j^{(m)}| \geq \sigma_{\mu\nu} \text{ for all } j \mu \neq i \nu \\ 0 & \text{if } |\vec{r}_i^{(n)} - \vec{r}_j^{(m)}| < \sigma_{\mu\nu} \text{ for one } j \mu \neq i \nu, \end{cases}$$

(2.17)

where $\sigma_{\mu\nu} = (\sigma_\mu + \sigma_\nu)/2$. This is the scheme we used in one of our studies in the next chapter.

To end this section, we point out that the MC method is a powerful tool to study equilibrium properties of a system. On the other hand, it does not follow the Newtonian laws for the particles. That’s why the molecular dynamics method (following section) is used when dynamic properties of a system are investigated.

2.2 Molecular dynamics

Molecular Dynamics simulation is a technique that computes the equilibrium and transport properties of a classical many-body system. It generates the microscopic information of the investigated system such as particles positions, velocities and so on. This information can be converted to the macroscopic properties of the system by statistical mechanics [30]. The motion of the constituent particles conforms to the laws of classical mechanics. In the following, we display our equations in a 3d system.

2.2.1 Equations of motion

Consider a system of $N$ particles which interact with the pair potential $u$. The forces imposed on particles derive from the potential energy (the system thus conserves the total energy). All the particles move in all directions of the system. The Lagrangian equation can be used to describe the motion of the particles. The equation (for the variable $q_k(t)$) is

$$\frac{d}{dt} \left( \frac{\partial L}{\partial \dot{q}_k} \right) - \left( \frac{\partial L}{\partial q_k} \right) = 0, \quad k = 1, \cdots 3N$$

(2.18)

where $q_k$ are the set of coordinates of all the particles ($k = 1, \ldots 3N$) and $\dot{q}_k$ are the time derivatives of $q_k$ (the momentum of the particles). The Lagrangian function $L(\{\dot{q}_k, q_k\})$ is
\[ \mathcal{L} = \mathcal{H} - \mathcal{V} \]  
(2.19)

with kinetic energy term \( \mathcal{H} = (1/2) \sum_{k=1}^{3N} \frac{\dot{q}_k^2}{2m} \) and potential energy term \( \mathcal{V} \). The total potential is

\[ \mathcal{V} = (1/2) \sum_{i=1}^{N} \sum_{j \neq i} u(r_{ij}) = \sum_{i=1}^{N-1} \sum_{j > i} u(r_{ij}), \]  
(2.20)

where we have \( \vec{r}_i = (x_i, y_i, z_i), \) \( x_1 = q_1, y_1 = q_2, z_1 = q_3, \) etc. The force on the particle \( i \) is

\[ \vec{F}_i = -\vec{\nabla}_{\vec{r}_i} \mathcal{V}. \]  
(2.21)

The force that atom \( j \) exerts on atom \( i \) is \( \vec{f}_{ij} = -\vec{\nabla}_{\vec{r}_i} u(r_{ij}). \) And for \( \vec{F}_i \) we have thus \( \vec{F}_i = \sum_{j \neq i} \vec{f}_{ij} = -\sum_{j \neq i} \vec{\nabla}_{\vec{r}_i} u(r_{ij}). \) According to Newton’s third law, \( \vec{f}_{ji} = -\vec{f}_{ij} \), meaning that each pair force needs only to be calculated once. If we take \( u \) to be the Lennard-Jones potential, then the pair force is

\[ \vec{f}_{ij,LJ} = \frac{48 \epsilon}{\sigma^2} \left[ \left( \frac{\sigma}{r_{ij}} \right)^{14} - \frac{1}{2} \left( \frac{\sigma}{r_{ij}} \right)^{8} \right] \vec{r}_{ij} \]  
(2.22)

The equation of motion for the particle \( i \) is

\[ m \ddot{\vec{r}}_i = \vec{F}_i = \sum_{j \neq i} \vec{f}_{ij} \]  
(2.23)

where the sum is over all \( N \) particles except \( i \) itself and \( m \) is the mass of the particle. These equations must usually be integrated numerically.

### 2.2.2 Finite difference methods

In molecular simulations, usually, the equations of motion can only be solved numerically because the force acting on particle \( i \) depends not only on its position \( \vec{r}_i \) but also on the positions of other particles \( \vec{r}_j (j \neq i) \), and in a non-linear manner (see for example eq.(2.22) for the Lennard-Jones pair force). An integrator is used to calculate the trajectory step by step with a finite time increment \( \delta t \). All the particles in a system evolve with time. What we want to know is the trajectory of the system during a period of time in the phase space. There are a lot of time integration algorithms to generate the MD trajectory. The most popular ones are the Gear predictor-corrector algorithm.
and the Verlet algorithm. We first show the Gear predictor-corrector algorithm as in
[30]. It proceeds in two steps. The first one is the prediction step. Suppose that at
time \( t \), the positions and velocities of the particles are known. Thus the accelerations
\( \vec{a}_i(t) = \vec{F}_i/m \) are known too. Suppose that we have also (a numerical estimate of)
the third time derivative of \( \vec{r}_i \) at \( t \), \( \vec{b}_i(t) \) and the fourth derivative \( \vec{c}_i(t) \). For a 5-value
predictor-corrector algorithm, the predicted positions and velocities, etc, at time \( t + \delta t \)
are given by a simple Taylor expansion
\[
\begin{align*}
\vec{r}_p^i(t + \delta t) &= \vec{r}_i(t) + \vec{v}_i(t)\delta t + \vec{a}_i(t)\delta t^2/2 + \vec{b}_i(t)\delta t^3/3! + \vec{c}_i(t)\delta t^4/4! \\
\vec{v}_p^i(t + \delta t) &= \vec{v}_i(t) + \vec{a}_i(t)\delta t + \vec{b}_i(t)\delta t^2/2 + \vec{c}_i(t)\delta t^3/3! \\
\vec{a}_p^i(t + \delta t) &= \vec{a}_i(t) + \vec{b}_i(t)\delta t + \vec{c}_i(t)\delta t^2/2 \\
\vec{b}_p^i(t + \delta t) &= \vec{b}_i(t) + \vec{c}_i(t)\delta t
\end{align*}
\]
(2.24)
The second step is the correction step. We calculate the accelerations \( \vec{a}_i(t + \delta t) \) from
the new (predicted) positions \( \{\vec{r}_p^i(t + \delta t)\} \), then the difference \( \Delta \vec{a}_i = \vec{a}_i(t + \delta t) - \vec{a}_i(t) \).
Using this, the corrected quantities are
\[
\begin{align*}
\vec{r}_c^i(t + \delta t) &= \vec{r}_p^i(t + \delta t) + G_0\Delta \vec{a}_i\delta t^2/2 \\
\vec{v}_c^i(t + \delta t) &= \vec{v}_p^i(t + \delta t) + G_1\Delta \vec{a}_i\delta t/2 \\
\vec{b}_c^i(t + \delta t) &= \vec{b}_p^i(t + \delta t) + G_3\Delta \vec{a}_i(3/\delta t) \\
\vec{c}_c^i(t + \delta t) &= \vec{c}_p^i(t + \delta t) + G_4\Delta \vec{a}_i(12/\delta t^2)
\end{align*}
\]
(2.25)
where, according to Gear, \( G_0 = 19/120, \quad G_1 = 3/4, \quad G_3 = 1/2 \) and \( G_4 = 1/12 \). The
corrected quantities allow to cumulate the statistics of the system, and constitute the
starting point of the next MD step.

A more direct and widely used method is the Verlet algorithm. We make sole
use of the equation of the motion, no need for the extra numerical coefficients. Its
velocity version (velocity Verlet) proceeds also by two steps. It has been shown to be
very stable, and is one of the most popular MD schemes. First, from the positions,
velocities , and the forces (thus the accelerations) of the particles at time \( t \), we compute
the positions at time \( t + \delta t \) and the velocities at time \( t + \delta t/2 \),
\[
\begin{align*}
\vec{r}_i(t + \delta t) &= \vec{r}_i(t) + \vec{v}_i(t)\delta t + \vec{a}_i(t)\delta t^2/2 \\
\vec{v}_i(t + \delta t/2) &= \vec{v}_i(t) + \vec{a}_i(t)\delta t/2
\end{align*}
\]
(2.26)
then, using the new positions \( \{ \vec{r}_i(t + \delta t) \} \), the forces (and the accelerations) at time \( t + \delta t \) are computed, and the velocities are advanced to \( t + \delta t \):

\[
\vec{v}_i(t + \delta t) = \vec{v}_i(t + \delta t/2) + \vec{a}_i(t + \delta t)\delta t/2.
\] (2.27)

The eqs.(2.26-2.27) constitute one step of the velocity Verlet algorithm. A comparison of the two methods has been made in [30]. It appears that if the forces only depend on the positions (not on the velocities) as in our systems, the velocity Verlet method can be preferred, since for relatively large \( \delta t \), it conserves better the total energy. Furthermore, it is simple to implement.

Whatever the integrator, our aim is to accumulate the statistic information of the system, and extract macroscopic properties. The results will depend on the ensemble used. The following gives some basic information about statistical ensembles that are relevant to our work.

### 2.3 Statistical ensembles

Statistical physics is used to study the thermodynamics of model materials. From the microscopic movement of particles composing the systems, it provides us with the relevant macroscopic properties. In the macroscopic world, the state of the system is characterized by some macroscopic quantities such as temperature, volume and pressure. In microscopic world, the classic statistical physics views systems as a mechanic system containing a very large number of particles whose movements conform to the Newtonian laws.

In statistical physics, ensembles are regarded as a large set of individual systems which have the same properties and structures in given macroscopic conditions. In our simulations, several ensembles are often encountered, including the microcanonical ensemble, the canonical ensemble and the isothermal-isobaric ensemble.

#### 2.3.1 Microcanonical ensemble

For the microcanonical ensemble, the number of the particles in the system, the volume and the total energy of the system are fixed to specific values (it is the NVE
ensemble). This is the natural ensemble for a standard MD simulation presented in
the section 2.2. The probability density for this ensemble is proportional to
\[ \delta (\mathcal{H} (\Gamma) - E) \] (2.28)
in this expression, \( \Gamma \) represents the phase space and \( \mathcal{H} (\Gamma) \) is the Hamiltonian. The
function describes the probability density of obtaining specific states of an N-particle
system in a space of volume \( V \) which have the desired energy \( E \). The partition function
is as follows
\[ Q_{NV E} = \sum_{\Gamma} \delta (\mathcal{H} (\Gamma) - E) \] (2.29)
its expression for a classical N-particles continuous system is
\[ Q_{NV E} = \frac{1}{N! h^{3N}} \int d\mathbf{r} d\mathbf{p} \delta (\mathcal{H} (\mathbf{r}, \mathbf{p}) - E) \] (2.30)
with \( h \) being the Planck constant. The correspondent thermodynamic potential is
(minus) entropy:
\[ -S = -k_B \ln Q_{NV E}. \] (2.31)
If, in the MD simulations, we were able to compute \( Q_{NV E} \), then according to eq.(2.31)
we can get access to the useful thermodynamics of our systems [33]. For example,
for the temperature, we have \( 1/T = \partial S/\partial E |_V \) and for the pressure, we have \( P/T = \partial S/\partial V |_E \). However, in an MD simulation, we do not compute the partition function
\( Q_{NV E} \), because we cannot possibly cover sufficiently the phase space in our runs. So
the entropy \( S \) is not computed. Nevertheless, many thermodynamic quantities can
be obtained by simply averaging their microscopic expressions during the run. For
example, the average kinetic energy is given by
\[ E_K = \langle \mathcal{H} \rangle_{MD}, \] (2.32)
leading to, according to the equipartition principle [32], the temperature
\[ T = \frac{2}{3 k_B} (E_K / N) \] (2.33)
Eq.(2.33) is the standard way of calculating the temperature of our systems.

In the NVE ensemble, \( E \) is fixed during the MD run. This means that when one
starts a simulation, one has the initial configuration, consisting of the initial positions
and initial velocities of the particles. If we set the initial velocities according to a
target temperature $T_0$ by using the Maxwell-Boltzmann distribution, the initial posi-
tions usually do not correspond to the equilibrium situation. They do fix the total
energy, giving $E = E_0$. During the MD run of equilibration, the system evolves to
the equilibrium state (maximum entropy) corresponding to given $(E, \rho)$, during which
there is exchange between the kinetic energy and potential energy. Consequently, the
equilibrium temperature will not be the target temperature $T_0$, but a different value.
This is a well-known difficulty of the standard MD simulation. It can be overcome
by rescaling the velocities regularly during the run (Andersen’s method), in order to
obtain an equilibrium configuration corresponding to the target temperature, meaning
that we run one kind of NVT MD for the equilibrium stage. Afterwards, NVE runs
can be carried out to study the system’s properties.

As the stress fluctuation formalism which we use is most directly case in the canonical (NVT) ensemble, we give in the following some basis of this ensemble.

### 2.3.2 Canonical ensemble

For the canonical ensemble, the number of particles in the system and the temper-
atture of the system are specified. The probability density is proportional to

$$\exp \left( -\mathcal{H}(\Gamma)/k_B T \right),$$

the partition function is

$$Q_{NVT} = \sum_{\Gamma} \exp \left( -\mathcal{H}(\Gamma)/k_B T \right),$$

the expression for an N-particle continuous system is

$$Q_{NVT} = \frac{1}{N!} \frac{1}{h^{3N}} \int d^N r \exp \left( -\mathcal{H}(\Gamma)/k_B T \right)$$

The corresponding thermodynamic potential is the Helmholtz free energy $F$

$$F = -k_B T \ln Q_{NVT}. $$

It is also convenient to define the configuration partition function, or the configuration
integral:

$$Z_{NVT} = \int d^N r \exp(-\mathcal{V}/k_B T).$$
\( Z_{NVT} \) is the relevant partition function, when we want to focus on the effects of particle interactions. As we mentioned earlier, the standard MC simulations (see section 2.1) are performed in the NVT ensemble. There is, of course, no question of calculating the partition function \( Q_{NVT} \), for the same reason as our not calculating \( Q_{NVE} \). However, we can compute plenty of physical quantities as MC averages, for example the energy, and its fluctuations. The latter is of course related to the constant-volume heat capacity of the system, as we shall see later.

Although the standard MD simulations are done in the NVE ensemble, nowadays it is common practice to perform them in the NVT ensemble. This can be done in several ways. One of them is the above-mentioned Andersen’s method. It however does not correspond to any deterministic dynamics, as opposed to the extended system Nose-Hoover method. In the latter, we consider the system in contact with a thermal reservoir. An extra degree of freedom is introduced, which represents the reservoir. Energy is allowed to flow between the system and the reservoir. The introduced degree of freedom is expressed as \( s \) and the conjugate momentum is \( p_s \). An extra potential energy is related to \( s \) as follows

\[
\mathcal{V}_s = (f + 1)k_B T \ln s \quad (2.39)
\]

Here \( f \) is the number of degrees of freedom and \( T \) is the specified temperature. The relevant kinetic energy is

\[
\mathcal{K}_s = \frac{1}{2} Q \dot{s}^2 = \frac{p_s^2}{2Q} \quad (2.40)
\]

Here \( Q \) is the thermal inertia parameter. It controls the rate of temperature fluctuations. The Lagrangian of the system is

\[
\mathcal{L}_s = \mathcal{K} + \mathcal{K}_s - \mathcal{V} - \mathcal{V}_s \quad (2.41)
\]

The equations of motion for the system can be derived

\[
\ddot{r} = \frac{f}{ms^2} - \frac{2\dot{s}\dot{r}}{s} \quad (2.42)
\]

and

\[
Q\ddot{s} = \sum_i m_i r_i^2 s - \frac{1}{s}(f + 1)k_B T \quad (2.43)
\]
The Hamiltonian $\mathcal{H}_s$ for the extended system is conserved

$$\mathcal{H}_s = \mathcal{H} + \mathcal{H}_s + V + V_s$$

(2.44)

and the density function for the extended system is microcanonical

$$\rho_{\text{NVE}}(r, p, s, p_s) = \frac{\delta(\mathcal{H}_s - E_s)}{\int drdpdsdp_s \delta(\mathcal{H}_s - E_s)}$$

(2.45)

This NVT scheme is used in our simulations. The NVT ensemble is important for our work. We shall show later how the stress fluctuation formalism is derived within this ensemble. Before this, we present another useful ensemble to our work.

### 2.3.3 Isothermal-isobaric ensemble

The isothermal-isobaric ensemble (NPT ensemble), is convenient for many studies because it fixes the temperature and the pressure of the system. This is similar to many experimental approaches. For this ensemble, the probability density is proportional to

$$\exp\left(-\frac{(\mathcal{H} + PV)}{k_B T}\right)$$

(2.46)

The partition function is

$$Q_{\text{NPT}} = \sum_{\Gamma} \sum_V \exp\left(-\frac{(\mathcal{H} + PV)}{k_B T}\right) = \sum_V \exp\left(-\frac{PV}{k_B T}\right) Q_{\text{NVT}}$$

(2.47)

The expression for an N-particle system is

$$Q_{\text{NPT}} = \frac{1}{N!} \frac{1}{h^{3N}} \frac{1}{V_0} \int dV \int d\mathbf{r} d\mathbf{p} \exp\left(-\frac{(\mathcal{H} + PV)}{k_B T}\right)$$

(2.48)

The appropriate thermodynamic potential is the Gibbs free energy $G$

$$G = -k_B T \ln Q_{\text{NPT}}$$

(2.49)

Again, we do not actually compute $G$ in our simulations. But we do calculate the averages and the fluctuations of physical quantities.

In order to perform MD simulations in fixed pressure $P$, we use extended system methods. Andersen proposed coupling the system to an external variable $V$, which is the volume of the simulation box. The coupling imitates the function of a piston on a real system. The kinetic energy for the piston is

$$\mathcal{K}_V = \frac{1}{2} Q \dot{V}^2$$

(2.50)
Here $Q$ is the mass of this piston. The extra potential energy for the system is

$$\mathcal{V}_V = PV$$  \hspace{1cm} (2.51)$$

Here $P$ is the specified pressure. The kinetic energy and potential energy related to the particles of the system are

$$\mathcal{V} = \mathcal{V} \left( V^\dagger s \right)$$  \hspace{1cm} (2.52)$$

$$\mathcal{K} = \frac{1}{2} m \sum_i v_i^2 = \frac{1}{2} m V^\dagger \sum_i \dot{s}_i^2$$  \hspace{1cm} (2.53)$$

Here $r = V^\dagger s$ and $v = V^\dagger \dot{s}$. The Lagrangian of the system is

$$\mathcal{L}_V = \mathcal{K} + \mathcal{K}_V - \mathcal{V} - \mathcal{V}_V$$  \hspace{1cm} (2.54)$$

The equations of motion for the system can be derived

$$\ddot{s} = \frac{f}{mV^\dagger} - \frac{2\dot{s}V}{3V}$$
$$\ddot{V} = \frac{P - \mathcal{P}}{Q}$$  \hspace{1cm} (2.55)$$

Here $f$ is the force and $\mathcal{P}$ is the pressure. They are calculated using unscaled coordinates and momenta. For the system, the Hamiltonian $\mathcal{H}_V$ is conserved.

$$\mathcal{H}_V = \mathcal{K} + \mathcal{K}_V + \mathcal{V} + \mathcal{V}_V.$$  \hspace{1cm} (2.56)$$

Eq. (2.55) corresponds actually to the dynamics of a constant NPH ensemble ($H = E + PV$ is the enthalpy). In order to carry out NPT runs, this scheme is coupled to the Nose-Hoover constant-temperature scheme shown earlier. This is the NPT-MD we used, within LAMMPS [29].

To perform isothermal-isobaric (NPT) MC simulations, the scheme of section 2.1 is also modified. We recall [30] that the configuration integral in this case is,

$$Z_{NPT} = \int dV \int d^N r \exp \left( -\beta (\mathcal{V} + PV) \right),$$  \hspace{1cm} (2.57)$$

with $V$ the volume variable. By performing the scaling $s = V^{-1/3} r$, we can rewrite $Z_{NPT}$ as

$$Z_{NPT} = \int dV V^N \int d^N s \exp \left( -\beta (\mathcal{V} + PV) \right)$$
$$= \int dV \int d^N s \exp \left( -\beta (\mathcal{V} + PV) + N \ln V \right).$$  \hspace{1cm} (2.58)$$
The new MC scheme then generates states consistent with the probability
\[ \rho_{NPT} \propto \exp \left( -\beta (\mathcal{V} + PV) + N \ln V \right), \tag{2.59} \]
by setting \( H = \mathcal{V} + PV - k_B T N \ln V \) and by considering the ensemble \( \{ s_1, \cdots, s_N, V \} \) as the \((3N+1)\) variables to which to apply random changes and the Metropolis algorithm
\[ P(m \rightarrow n) = \min \left( 1, \exp \left( -\beta (H_n - H_m) \right) \right) \tag{2.60} \]
where \( H_m = \mathcal{V}_m + PV_m - k_B T N \ln V_m \) and similarly for \( H_n \). Eq.(2.60) is used to our investigation of HS systems under constant-NPT condition (see next chapter).

### 2.4 Simple averages and fluctuations

Our simulations lead us to calculate various physical quantities. Some are simple averages, other are the fluctuations of relevant quantities which are related to the response functions of our systems. As explained in [30], for the same thermodynamic state, simple averages are independent of the ensemble used, whereas the fluctuations depend on the ensemble. For example, if we take a system defined by its density \( \rho = N/V \) and its temperature \( T \) in the NVT ensemble, at equilibrium, the energy is given by
\[ E = \langle \mathcal{H} \rangle_{NVT}. \tag{2.61} \]
Equivalently, we can run NVE simulations, with the same \( E \), and we shall obtain the (average)equilibrium temperature by eq.(2.33), which is the same as the one of our NVT ensemble. This idea also applies to the calculation of the pressure \( P \). Running (equilibrium) NVT simulations gives the pressure \( \bar{P} \). If we run NPT simulations, then we shall obtain average volume \( \bar{V} \). Again, if we have \( \bar{P} = P \), then \( \bar{V} = V \). More precisely, here are some common simple averages
\[
\begin{align*}
E & = \langle \mathcal{H} \rangle \\
E_K & = \langle \mathcal{H} \rangle = (1/2) \left\langle \sum_i m_i v_i^2 \right\rangle \\
E_P & = \langle \mathcal{V} \rangle = \left\langle \sum_{i=1}^{N-1} \sum_{j>i} u(r_{ij}) \right\rangle \\
T & = (2/3)E_K/Nk_B
\end{align*}
\tag{2.62} \]
for total energy, kinetic energy, potential energy and temperature respectively.

The pressure $P$ is an important quantity in our studies. Its expression is well-known for pairwise potentials. To show this, we start from the definition $P = -(\partial F/\partial V)_T$. As $F = -k_B T \ln Q_{NVT}$, we have

$$P = \frac{k_B T}{Q_{NVT}} \left( \frac{\partial Q_{NVT}}{\partial V} \right)_T$$

(2.63)

From eq.(2.36) and eq.(2.38)we have

$$Q_{NVT} = A(T)Z_{NVT} = A(T) \int d^N \mathbf{r} \exp(-\mathcal{V}/k_B T).$$

(2.64)

where $A(T) = (2\pi m k_B T / h^3)^N / N!$ is independent of the volume $V$. By the scaling method $\mathbf{r}' = V^{1/3} \mathbf{s}$, we can rewrite $Z_{NVT}$ as

$$Z_{NVT} = V^N \int d^N \mathbf{s} \exp(-\mathcal{V}/k_B T).$$

(2.65)

As $\mathcal{V} = \sum_{i=1}^{N-1} \sum_{j>i} u(r_{ij}) = \sum_{i=1}^{N-1} \sum_{j>i} u(V^{1/3}s_{ij})$, we have

$$(\partial \mathcal{V}/\partial V)_T = \sum_{i=1}^{N-1} \sum_{j>i} u'(r_{ij})s_{ij}(1/3)V^{-2/3} = (1/3V) \sum_{i=1}^{N-1} \sum_{j>i} u'(r_{ij})r_{ij}$$

(2.66)

Using eq.(2.63) to eq.(2.66), we obtain finally,

$$P = \rho k_B T - (1/3V) \left\{ \sum_{i=1}^{N-1} \sum_{j>i} u'(r_{ij})r_{ij} \right\}.$$  

(2.67)

This is the famous virial expression for the pressure. The first term is the ideal gas term, $P_{id}$, the second one is the “excess” term $P_{ex}$, due to the interactions between the particles.

Another common simple average we compute is the radial distribution function $g(r)$, which characterizes the local structure in our systems. The function $\rho g(r)$ represents the probability density of finding a particle at $r$, given that another is at the origin. Furthermore, in an isotropic system, $g(r)$ is related to the thermodynamic quantities of the system,

$$E_p(\text{isotropic})/N = 2\pi \rho \int dr r^2 g(r)u(r)$$

(2.68)

$$P(\text{isotropic}) = \rho k_B T - (2\pi/3)\rho^2 \int dr g(r)u'(r)r^2.$$  

(2.69)
This function will be studied in detail in the next chapter.

Now we turn to the fluctuations. The fluctuations depend sensibly on the ensemble. Take for example the energy $E$. In an NVE ensemble, $E$ does not fluctuate, i.e. $\Delta E \equiv \langle (H - E)^2 \rangle = 0$. On the other hand, we have [30]

$$\Delta E_{NVT} = k_B T^2 C_V > 0,$$

where $C_V = (\partial E / \partial T)_V$ is the specific heat capacity of the system. So the fluctuations of $E$ in the NVT ensemble is a way of calculating the constant-volume specific heat capacity. (Functions like $C_V$ are often called “response functions”). Another example, more related to our work, is the fluctuations of the volume $V$. Obviously, in an NVT ensemble, $\Delta V = 0$, whereas in an NPT ensemble, we have [30]

$$\Delta V_{NPT} \equiv \langle (V - \bar{V})^2 \rangle = \bar{V} k_B T \kappa_T > 0,$$

where $\kappa_T = -V^{-1} (\partial V / \partial P)_T$ is the isothermal compressibility of the system. Using the bulk modulus $K = 1 / \kappa_T$, eq.(2.71) can be rewritten as

$$K = \frac{\bar{V} k_B T}{\Delta V_{NPT}}.$$  (2.72)

We see that eq.(2.72) provides a means of computing the bulk modulus. Thus this relation is very useful to our work, as shown in the next chapter.

### 2.5 Calculation methods of elastic properties

Here we present the basic formulas of the calculation of elastic properties by the stress fluctuation formalism. We are in general situations of $P \neq 0$, and $T \neq 0$. The simulations are equilibrium simulations, i.e. our systems are never actually deformed or submitted to shear stress. On the other hand, we are restricted to pairwise potentials.

#### 2.5.1 Elasticity of solids under pressure

The formalism is most conveniently derived in the canonical ensemble.

We follow the notations in [34]. Given a system $(N, V, T)$, let $\vec{X}$ be the initial configuration, $\vec{x}$ the final configuration. The displacement gradient $u_{\alpha\beta}$ is

$$x_\alpha - X_\alpha = u_\alpha(\vec{X});\ u_{\alpha\beta} = \partial u_\alpha / \partial X_\beta$$  (2.73)
The conventional strain tensor is
\[ \epsilon_{\alpha\beta} = \frac{1}{2} (u_{\alpha\beta} + u_{\beta\alpha}) \]  
(2.74)

The distance \( R_{ij} = |\vec{R}_i - \vec{R}_j| \) changes to \( r_{ij} \) according to
\[ r_{ij}^2 = R_{ij}(1 + \epsilon^T)(1 + \epsilon)R_{ij} \]
(2.75)
where
\[ \eta = \frac{1}{2} (\epsilon + \epsilon^T + \epsilon^T \epsilon) \]
(2.76)
is the *Lagrangian strain*. The free energy per unit of (the undeformed) volume \( f = F/V_0 \), can be expanded in powers of \( \eta \):
\[ f(\vec{x}, T) = f(\vec{X}, \eta_{\alpha\beta}, T) = f(\vec{X}, 0, T) + C_{\alpha\beta} \eta_{\alpha\beta} + \frac{1}{2} C_{\alpha\beta\chi\kappa} \eta_{\alpha\beta} \eta_{\chi\kappa} + \cdots \]  
(2.77)
where \( C_{\alpha\beta\chi\kappa} \) are the elastic constants.

The stress \( \sigma_{\alpha\beta} \) is related to the 1st derivative of \( f \) by
\[ \sigma_{\alpha\beta}(\vec{x}) = \det(h)^{-1} h_{\alpha\chi} \frac{\partial f}{\partial \eta_{\chi\kappa}}(\vec{x}) h_{\kappa\beta} \]  
(2.78)
where \( h = I + \epsilon \). And the stress-strain relation is:
\[ \sigma_{\alpha\beta} = B_{\alpha\beta\chi\kappa} \epsilon_{\chi\kappa} \]  
(2.79)
where \( B_{\alpha\beta\chi\kappa} \) are the Birch coefficients [34]. They are the elastic constants implied in the Hooke’s law eq. (2.79).

In case \( \sigma_{\alpha\beta}(\vec{X}) = -P \delta_{\alpha\beta} \) (isotropic initial stress), we have
\[ B_{\alpha\beta\chi\kappa} = C_{\alpha\beta\chi\kappa} + P(\delta_{\alpha\beta}\delta_{\chi\kappa} - \gamma_{\alpha\chi} \gamma_{\beta\kappa} - \delta_{\alpha\kappa} \delta_{\beta\chi}) \]  
(2.80)
Obviously, if \( P = 0 \), we have \( B = C \). This is often assumed in many textbooks on the elasticity of the solids.

By the Voigt notation, we write \( 1 = xx, 2 = yy, 3 = zz, 4 = yz \) (or \( yz \)), \( 5 = zx \) (or \( zx \)), \( 6 = xy \) (or \( yx \)). Thus, the strain tensor reads,
\[ \sigma = \begin{pmatrix} \sigma_1 & \sigma_6 & \sigma_5 \\ \sigma_6 & \sigma_2 & \sigma_4 \\ \sigma_5 & \sigma_4 & \sigma_3 \end{pmatrix} \]  
(2.81)
And the stress tensor reads, by convention

\[
\varepsilon = \begin{pmatrix}
\varepsilon_1 & \varepsilon_6/2 & \varepsilon_5/2 \\
\varepsilon_6/2 & \varepsilon_2 & \varepsilon_4/2 \\
\varepsilon_5/2 & \varepsilon_4/2 & \varepsilon_3 \\
\end{pmatrix}
\]  
(2.82)

**Special case one: 3-dimensional Cubic lattice**

For a cubic lattice, the \( B \) tensor is written as

\[
B = \begin{pmatrix}
B_{11} & B_{12} & B_{12} & 0 & 0 & 0 \\
B_{12} & B_{11} & B_{12} & 0 & 0 & 0 \\
B_{12} & B_{12} & B_{11} & 0 & 0 & 0 \\
0 & 0 & 0 & B_{44} & 0 & 0 \\
0 & 0 & 0 & 0 & B_{44} & 0 \\
0 & 0 & 0 & 0 & 0 & B_{44} \\
\end{pmatrix},
\]  
(2.83)

where \( B_{44} = G \) is the shear modulus. The bulk modulus is given by \( K = (B_{11} + 2B_{12})/3 \).

The matrix inverse to \( B \) (the compliance tensor \( S \)) is thus (for cubic symmetry):

\[
S = B^{-1} = \begin{pmatrix}
b_1 & b_2 & b_2 & 0 & 0 & 0 \\
b_2 & b_1 & b_2 & 0 & 0 & 0 \\
b_2 & b_2 & b_1 & 0 & 0 & 0 \\
0 & 0 & 0 & G^{-1} & 0 & 0 \\
0 & 0 & 0 & 0 & G^{-1} & 0 \\
0 & 0 & 0 & 0 & 0 & G^{-1} \\
\end{pmatrix}
\]  
(2.84)

where

\[
b_1 = \frac{B_{11} + B_{12}}{(B_{11} - B_{12})(B_{11} + 2B_{12})},
\]

\[
b_2 = -\frac{B_{12}}{(B_{11} - B_{12})(B_{11} + 2B_{12})}.
\]  
(2.85)

Thus the Young modulus is

\[
E = \frac{1}{b_1} = \frac{(B_{11} - B_{12})(B_{11} + 2B_{12})}{B_{11} + B_{12}},
\]  
(2.86)

and the Poisson ratio is given by

\[
\nu = \frac{b_2}{b_1} = \frac{B_{12}}{B_{11} + B_{12}}.
\]  
(2.87)
General d-dimensional cubic (square) lattice (d=2, 3)

For a 2-dimensional system possessing 3 elastic constants (for example the square lattice), the previous equations are easily rewritten accordingly, by suppressing the z-component. We have thus $B_{11}$, $B_{12}$ and $B_{66}$ as independent Hooke’s constants. It is possible to summarise the general $d$-dimensional situations in a unique formulation. Indeed, while the relation

$$G = B_{66} = C_{66} - P$$  \hspace{1cm} (2.88)

is $d$-independent, the relation

$$K = \frac{1}{d} [B_{11} + (d - 1)B_{12}] = \frac{1}{d} [C_{11} + (d - 1)C_{12} + (d - 2)P]$$  \hspace{1cm} (2.89)

depends on $d$. Of course, the compliance tensor is also affected. Thus, we have

\begin{align*}
b_1 &= \frac{B_{11} + (d - 2)B_{12}}{B_{11}^2 - (d - 2)B_{11}B_{12} - (d - 1)B_{12}^2} \hspace{1cm} (2.90) \\
b_2 &= -\frac{B_{12}^2 - (d - 2)B_{11}B_{12} - (d - 1)B_{12}^2}{B_{11} + (d - 2)B_{12}}
\end{align*}

with the Young modulus given by

$$E = \frac{1}{b_1} = \frac{B_{11}^2 - (d - 2)B_{11}B_{12} - (d - 1)B_{12}^2}{B_{11} + (d - 2)B_{12}},$$  \hspace{1cm} (2.91)

and the Poisson ratio given by

$$\nu = -\frac{b_2}{b_1} = \frac{B_{12}}{B_{11} + (d - 2)B_{12}}.$$  \hspace{1cm} (2.92)

While eqs.(2.88,2.89,2.91) allow to obtain $K$, $G$ and $E$ from the $B$ constants, it is also interesting to discuss the inverse problem: knowing these quantities from experimental measures, how can we deduce the $B$s? The answer is quite straightforward, by inverting these equations, leading to

\begin{align*}
B_{11} &= d.K \frac{d.K + (d - 2)E}{d^2.K - E} \\
B_{12} &= d.K \frac{d.K - E}{d^2.K - E} \hspace{1cm} (2.93) \\
B_{66} &= G.
\end{align*}
**Isotropic d-dimensional systems**

For an isotropic material, the Lamé coefficients \( \lambda \) and \( \mu \) are usually introduced. They are related to the \( C \) constants by

\[
C_{\alpha\beta\chi\kappa} = \lambda \delta_{\alpha\beta} \delta_{\chi\kappa} + \mu (\delta_{\alpha\chi} \delta_{\beta\kappa} + \delta_{\alpha\kappa} \delta_{\beta\chi}).
\]

(2.94)

Thus

\[
B_{\alpha\beta\chi\kappa} = (\lambda + P) \delta_{\alpha\beta} \delta_{\chi\kappa} + (\mu - P)(\delta_{\alpha\chi} \delta_{\beta\kappa} + \delta_{\alpha\kappa} \delta_{\beta\chi})
\]

(2.95)
yielding \( B_{11} = \lambda + 2\mu - P \), \( B_{12} = \lambda + P \), and \( B_{66} = \mu - P = G \). These are \( d \)-independent relations.

In terms of \( \lambda \) and \( \mu \), the bulk modulus \( K = -V \frac{\partial P}{\partial V} \) is given by

\[
K = \frac{1}{d}(B_{11} + (d - 1)B_{12}) = \lambda + 2\mu/d + (d - 2)P/d,
\]

(2.96)

the shear modulus \( G \) is

\[
G = \mu - P.
\]

(2.97)

Inversely, we have \( \lambda = K - 2G/d - P \), \( \mu = G + P \). Thus, from eq.(2.95), we can express the relevant \( B \)s in terms of \( K \) and \( G \):

\[
\begin{align*}
B_{11} &= K + 2(d - 1)G/d \\
B_{12} &= K - 2G/d \\
B_{66} &= G
\end{align*}
\]

(2.98)

Of course, here we have \( (B_{11} - B_{12})/2 = G = B_{66} \). In this case, the Young modulus can be obtained from eq.(2.93) and eq.(2.98), leading to

\[
E = \frac{2d^2K \cdot G}{d(d - 1)K + 2G} = 2(\mu - P) \frac{d \cdot \lambda + 2\mu + (d - 2)P}{(d - 1)\lambda + 2\mu + (d - 3)P},
\]

(2.99)

and the Poisson ratio, from eq.(2.92), gives

\[
\nu = \frac{d \cdot K - 2G}{d(d - 1)K + 2G} = \frac{\lambda + P}{(d - 1)\lambda + 2\mu + (d - 3)P}.
\]

(2.100)
Two-dimensional triangular lattice

For a triangular lattice (2d), $B$ is a $3 \times 3$ tensor. Only two independent $B$ constants exist. They are $B_{11} = B_{22}$ and $B_{12} = B_{21}$. For the shear modulus $B_{66}$, we have $B_{66} = (B_{11} - B_{12})/2$. The other elements are zero. We see that the situation (concerning the elasticity) is similar to the case of a two-dimensional isotropic solid (glass).

2.5.2 Stress fluctuation formalism for crystalline solids

Within the stress fluctuation formalism [8], we have, supposing the Hamiltonian $H = \sum_i p_i^2/(2m) + \sum_{i,j} u(r_{ij})$, the first term being the kinetic part, the second being the potential energy part, (where $\sum_{i<j}$ is a short-handed notation for $\sum_i \sum_{j>i}$), and using eqs.(2.75) - (2.80), the elastic constants $C$ expressed as the sum of three terms:

$$C_{\alpha\beta\chi\kappa} = C^K_{\alpha\beta\chi\kappa} + C^B_{\alpha\beta\chi\kappa} - C^F_{\alpha\beta\chi\kappa} \tag{2.101}$$

where $C^K$ is the kinetic part

$$C^K_{\alpha\beta\chi\kappa} = 2k_B T \rho (\delta_{\alpha\kappa} \delta_{\beta\chi} + \delta_{\alpha\chi} \delta_{\beta\kappa}) \tag{2.102}$$

with the density $\rho$. The Born part is well-known:

$$C^B_{\alpha\beta\chi\kappa} = \frac{1}{V} \sum_{i<j} \left( \frac{u'(r_{ij}) - u'(r_{ij})}{r_{ij}} \right) \frac{r_{ij}^\alpha r_{ij}^\beta r_{ij}^\chi r_{ij}^\kappa}{r_{ij}^2} \tag{2.103}$$

And the fluctuation part is

$$C^F_{\alpha\beta\chi\kappa} = \frac{V}{k_B T} \left[ \langle \hat{\sigma}_{\alpha\beta} \hat{\sigma}_{\chi\kappa} \rangle - \langle \hat{\sigma}_{\alpha\beta} \rangle \langle \hat{\sigma}_{\chi\kappa} \rangle \right] \tag{2.104}$$

with the stress tensor element $\hat{\sigma}_{\alpha\beta}$ given by, for pairwise potentials:

$$\hat{\sigma}_{\alpha\beta} = \frac{1}{V} \left( \sum_{i<j} \frac{u'(r_{ij}) r_{ij}^\alpha r_{ij}^\beta}{r_{ij}} - \sum_i p_i^\alpha p_i^\beta \right) \tag{2.105}$$

We note that in eq.(2.104), the fluctuation term is defined as the opposite of that in [8]. This choice is made for numerical convenience (see next chapter). We can split $C^F$
into “kinetic” and “configurational” parts:

\[ C_{\alpha\beta\chi\kappa}^{FK} = \frac{1}{V k_B T} \left[ \sum_i \sum_j \left\langle \frac{p_i^{\alpha} p_i^{\beta} p_j^{\chi} p_j^{\kappa}}{m^2} \right\rangle \right. \]

\[ \left. + \sum_i \left\langle \frac{p_i^{\alpha}}{m^2} \right\rangle \sum_j \left\langle \frac{p_j^{\chi}}{m^2} \right\rangle \right] \tag{2.106} \]

\[ = \rho k_B T (\delta_{\alpha\chi} \delta_{\beta\kappa} + \delta_{\alpha\kappa} \delta_{\beta\chi}) \]

and

\[ C_{\alpha\beta\chi\kappa}^{FC} = \frac{1}{V k_B T} \left[ \sum_{i<j} \sum_{k<l} \left\langle \frac{u'(r_{ij}) u'(r_{kl}) r_{ij}^{\alpha} r_{ij}^{\beta} r_{kl}^{\chi} r_{kl}^{\kappa}}{r_{ij} r_{kl}} \right\rangle \right. \]

\[ \left. + \sum_{i<j} \left\langle \frac{u'(r_{ij}) r_{ij}^{\alpha} r_{ij}^{\beta}}{r_{ij}} \right\rangle \sum_{i<j} \left\langle \frac{u'(r_{ij}) r_{ij}^{\chi} r_{ij}^{\kappa}}{r_{ij}} \right\rangle \right] \tag{2.107} \]

As we can see, the 1st term in formula (2.107) implies 3-particle and 4-particle distribution functions. We have now

\[ C = C^B - C^{FK} + C^{KK} \]

with \( C^{KK} = C^K - C^{FK} \), i.e.

\[ C_{\alpha\beta\chi\kappa}^{KK} = \rho k_B T (\delta_{\alpha\chi} \delta_{\beta\kappa} + \delta_{\alpha\kappa} \delta_{\beta\chi}) \tag{2.108} \]

To split completely the “ideal gas” term and the “excess” term (due to the interactions), we can rewrite \( C \) as

\[ C_{\alpha\beta\chi\kappa} = C_{\alpha\beta\chi\kappa}^{B} - C_{\alpha\beta\chi\kappa}^{FC} + C_{\alpha\beta\chi\kappa}^{KK} \tag{2.109} \]

\[ C_{\alpha\beta\chi\kappa}^{id} = C_{\alpha\beta\chi\kappa}^{KK} = \rho k_B T (\delta_{\alpha\chi} \delta_{\beta\kappa} + \delta_{\alpha\kappa} \delta_{\beta\chi}) \tag{2.110} \]

\[ C_{\alpha\beta\chi\kappa}^{ex} = C_{\alpha\beta\chi\kappa}^{B} - C_{\alpha\beta\chi\kappa}^{FC} \tag{2.111} \]

As we shall see later, in most of our (dense) systems, the contribution of the ideal term (proportional to \( T \)) is negligible. The Born term is usually more important than the fluctuation term, with the latter not at all neglectable, except for simple crystals at \( T = 0 \).

As we always compute the bulk modulus \( K \) and the shear modulus \( G \) in our systems, we give them specific expressions in the following.

Recalling that \( K = -V (\partial P/\partial V)_T = [C_{11} + (d - 1)C_{12} + (d - 2)P] / d \), we can show that after some algebra, the bulk modulus \( K \) can be expressed as

\[ K = P + \frac{\langle \chi \rangle}{V} - \langle \delta P^2 \rangle (\frac{V}{k_B T}) \tag{2.112} \]
here the “hypervirial function” [30], also referred to as the “Born-Lamé coefficient” [35] is

\[ \langle \chi \rangle = \frac{1}{d^2} \left\langle \sum_{i<j} r_{ij} d \left( r_{ij} u'(r_{ij}) \right) \right\rangle \tag{2.113} \]

and we have \( P = P_{id} + P_{ex} \) with \( P_{id} = \rho k_B T \), and

\[ P_{ex} = \langle P_{ex} \rangle = -\frac{1}{d} \cdot V \left\langle \sum_{i<j} r_{ij} u'(r_{ij}) \right\rangle \tag{2.114} \]

is the “virial equation”. Thus \( \delta P_{ex} = P_{ex} - P_{ex} \). We see that eq.(2.112) is an isotropic expression, as expected, for the bulk modulus.

For the shear modulus \( G \), we have

\[ G = C_{66} - P = C_{66}^B - C_{66}^{FC} + C_{66}^{KK} - P \]
\[ = C_{66}^B - C_{66}^{FC} - P_{ex} \tag{2.115} \]

In general, this is not an isotropic property, depending on the symmetry of the crystal. We shall examine it in more details later.

Coming back to \( K \), and following our recent works, it is convenient to write

\[ K = P_{id} + \eta_{A,ex} - \eta_{F,ex} \tag{2.116} \]

where \( \eta_{A,ex} \) is the elastic bulk modulus corresponding to the excess affine elasticity of the system. More precisely, we have

\[ \eta_{A,ex} = P_{ex} + \frac{\langle \chi \rangle}{V} \tag{2.117} \]

and

\[ \eta_{F,ex} = \langle \delta P_{ex}^2 \rangle \frac{V}{k_B T} \tag{2.118} \]

For pairwise potentials and \( T > 0 \), obviously, \( \langle \chi \rangle/V \) and \( P_{ex} \) (thus the affine parts) can be expressed from the radial distribution function (RDF) \( g(r) \) and the pair potential \( u(r) \), i.e.

\[ \frac{\langle \chi \rangle}{V} = \frac{\rho^2}{2d^2} \int dr \ S_d g(r) r \frac{d(r u'(r))}{dr} \tag{2.119} \]

where \( S_d \) is the surface of a \( d \)-sphere \( (S_3 = 4\pi r^2, S_2 = 2\pi r) \). and

\[ P_{ex} = -\frac{\rho^2}{2d} \int dr \ S_d g(r) r \ u'(r) \tag{2.120} \]
In a crystal at \( T = 0 \), eq.(2.113) can be simply written as
\[
\frac{\langle \chi \rangle}{V} = \frac{\rho}{2d^2} \sum_k s_k r_k \frac{d}{dr_k} (r_k u'(r_k))
\] (2.121)
where \( r_k \) are the coordination shells, and \( s_k \) are the coordination numbers for each shell. The summation terminates when \( r_k \geq r_c \), the cutoff distance of the pair potential \( u(r) \).

For example, the triangular lattice with lattice constant \( a \), we have \( r_k = (a, \sqrt{3}a, 2a, ...) \) and \( s_k = (6, 6, 6, ...) \). The lattice constant, related to the density by \( \rho = 2/\sqrt{3}a^2 \), is of course determined by the fixed pressure \( P \), where for \( T = 0, P = P_{ex} \), from eq. (2.114) is
\[
P_{ex} = -\frac{\rho}{2d} \sum_k s_k r_k u'(r_k)
\] (2.122)
In this way, we can compute \( \eta_{A,ex} \) at \( T = 0 \). However, \( \eta_{F,ex} \) cannot be computed using (2.118). It will be estimated from extrapolation of low temperature results.

Now some details about \( G \). The computation of \( G \) in a cubic crystal follows eq.(2.115), with the symmetry \( C_{66} = C_{55} = C_{44} \) (numerically, it is possible to average these three elements). Next we turn to the case of a 2d triangular lattice. In such a symmetry, we also have a second formula for the shear modulus, denoted by \( G_2 \),
\[
G_2 = \frac{(C_{11} - C_{12})}{2} - P
\]
\[
= \frac{(C_{11}^B - C_{12}^B - C_{11}^{FC} + C_{12}^{FC})}{2} - P_{ex}
\] (2.123)
By denoting \( G_1 \) the result given by eq.(2.115), we can write the averaged \( G \) as \( G_{2d} = (G_1 + G_2)/2 \). This averaging is applied to the study of low temperature monodisperse 2d LJ system, forming a triangular lattice [36].

2.5.3 Elastic moduli of the glassy state

The glassy phase is an isotropic solid. Its elastic moduli \( K \) and \( G \) fully characterize its elasticity. For \( K \), the equation is given by eq.(2.116), i.e. \( K = P_{id} + \eta_{A,ex} - \eta_{F,ex} \). As for \( G \), starting from eq.(2.115), we can express it similarly as
\[
G = \mu_{A,ex} - \mu_{F,ex}
\] (2.124)
with
\[
\mu_{A,ex} = \langle C_{66}^{B,ang} \rangle - P_{ex} = \frac{d}{d+2} \left( \frac{\langle \chi \rangle}{V} - P_{ex} \right) = \frac{d}{d+2} (\eta_{A,ex} - 2P_{ex}),
\] (2.125)
with $\langle C_{66}^B \rangle_{\text{ang}}$ being the angular average of $C_{66}^B$ in the $d$-space, and

$$
\mu_{F, \text{ex}} = C_{66}^{FC} = \frac{V}{k_B T} \langle \delta \hat{\sigma}_{xy, \text{ex}}^2 \rangle.
$$

(2.126)

We see from eq.(2.125) that $\eta_{A, \text{ex}}$ and $\mu_{A, \text{ex}}$ are related. This is not the case for the non-affine part of $K$ and $G$, $\eta_{F, \text{ex}}$ and $\mu_{F, \text{ex}}$. These quantities, involving three- and four-particles correlations, must be computed from the stress fluctuations. As the case of $\eta_{F, \text{ex}}$, $\mu_{F, \text{ex}}$ given by eq.(2.126) can only be computed for $T > 0$. So its value at $T = 0$ will be an extrapolation of the results for $T > 0$. We shall see that, depending on the complexity of the crystal, it is not necessarily zero.

Our glass formers are often binary mixtures. We adopt here eqs.(2.119) and (2.120) to the case of a binary mixture:

$$
\frac{\langle \chi \rangle}{V} = \sum_{a=1}^{2} \sum_{b=1}^{2} \frac{\rho_a \rho_b}{2d^2} \int dr S_d(r) g_{ab}(r) \frac{d(r u'_{ab}(r))}{dr} \tag{2.127}
$$

where $a$ and $b$ run over the species, and

$$
P_{\text{ex}} = -\sum_{a=1}^{2} \sum_{b=1}^{2} \frac{\rho_a \rho_b}{2d} \int dr S_d(r) g_{ab}(r) \frac{d(r u'_{ab}(r))}{dr} \tag{2.128}
$$

As for the equation of $\hat{\sigma}_{\alpha \beta}$, eq.(2.105), its extension to mixtures is straightforward, since the sums run over each particle of the system.

### 2.5.4 Elastic moduli of the liquid state

In the liquid state, there is no well-defined displacement field for the particles. What we know is $K = -(1/V)(\partial P/\partial V)_T > 0$ and $G = 0$. It seems difficult to transpose previous results to liquids, since they are based on explicit displacement fields. For the compression modulus, Rowlinson [9] demonstrated eq.(2.112) for $K$ via a discussion of the fluctuations of $P_{\text{ex}}$. The system was put in a cubic box of length $L$. One writes for the particle $i$

$$
r_{ia} = s_{ia} L \tag{2.129}
$$

where $\alpha = x, y, z \ (d = 3)$, $(x, y$ for $d = 2)$, the reduced coordinates $0 \leq s_{ia} \leq 1$. As the pressure $P = -(\partial F/\partial V)_T = P_{td} + P_{\text{ex}}$, with $V = L^d$, $F$ the free energy, one can show that $P_{td} = \rho k_B T$, and

$$
P_{\text{ex}} = \frac{k_B T}{dL^{d-1}Z} \left( \frac{\partial Z}{\partial L} \right)_T \tag{2.130}
$$
with \( Z \) the canonical configuration partition function, defined by eq.(2.38). This gives the virial expression of \( P \) (see eq. 2.67). As the bulk modulus \( K = -V(\partial P/\partial V)_T = \rho k_B T - V(\partial P_{ex}/\partial V)_T \), we need to differentiate once more the second member of eq.(2.130) with respect to \( V \) (thus \( L \)). Using again the “trick” eq.(2.129) and after some algebra, eq.(2.112) can be proven. We note that the proof can be extended to a rectangular box with fixed aspect ratios.

In a liquid, obviously, \( G = 0 \). The fluctuation expression (2.124) is indeed consistent with this fact. As Zwanzig showed [37], we have

\[
\langle \hat{\sigma}^2_{xy,ex} \rangle = \langle \hat{\sigma}^2_{xy,ex} \rangle - \langle \hat{\sigma}_{xy,ex} \rangle^2
\]

\[
= \frac{1}{V^2} \left( \sum_i x_i F_i^y \sum_k x_k F_k^y \right)
\]

\[
= -\frac{k_B T}{V^2} \left( \sum_i \partial_i \left( \sum_k x_k F_k^y \right) / \partial y_i \right)
\]

\[
= -\frac{k_B T}{V^2} \left( \sum_i x_i \partial \left( \sum_k \sum_{l>k} x_{kl} F_{kl}^y \right) / \partial y_i \right)
\]

\[
= \frac{k_B T}{V^2} \left( \sum_i \sum_{j>i} x^2_{ij} \partial^2 u(r_{ij}) / \partial y_{ij}^2 \right)
\]

\[
= \frac{k_B T}{V^2} \left( \sum_i \sum_{j>i} x^2_{ij} \left[ \frac{u'(r_{ij})}{r_{ij}} + \frac{u'' - u'}{r_{ij}^2} \frac{y_{ij}^2}{r_{ij}} \right] \right).
\]

Note that in eq.(2.131), we assumed \( \langle \hat{\sigma}_{xy,ex} \rangle = 0 \). This is true in a static liquid, and is well verified in simulations. After spherical average (in the \( d \)-space), we obtain

\[
\langle \delta \hat{\sigma}^2_{xy,ex} \rangle = \frac{k_B T}{dV^2} \left( \sum_i \sum_{j>i} \left[ r_{ij} u' + u'' \frac{r_{ij}^2}{d+2} \right] \right)
\]

\[
= \frac{k_B T}{d(d+2)V^2} \left( \sum_i \sum_{j>i} \left[ (d+1)r_{ij} u' + u'' r_{ij}^2 \right] \right)
\]

\[
= \frac{d k_B T}{(d+2)V} \left[ -P_{ex} + \langle \chi \rangle / V \right]
\]

Putting eq.(2.132) in eq.(2.124). One shows indeed \( G = 0 \) for liquids. We note that using the radial distribution function \( g(r) \), we can also write, for a one-component system

\[
\beta V \langle \delta \hat{\sigma}^2_{xy} \rangle = \frac{\rho^2}{2d(d+2)} \int dr S_d g(r) [(d+1)r u'(r) + r^2 u''(r)]
\]
and for mixtures

\[
\beta V \langle \delta \hat{\sigma}_{xy}^2 \rangle = \sum_a \sum_b \frac{\rho_a \rho_b}{2d(d+2)} \int dr \ S_d g_{ab}(r) \left[ (d+1)ru'_{ab}(r) + r^2 u''_{ab}(r) \right]
\]

(2.134)

where \( a \) and \( b \) run over the species. Thus, in liquids, the fluctuation term does not involve higher order (3 or 4) distribution functions.

### 2.5.5 Additional theoretical aspects

Before closing the section, we point out that the stress fluctuation formulas in subsections 2.5.2 - 2.5.4 are obtained (and only valid) within the NVT ensemble, more precisely, the \( NV \gamma T \) ensemble, with \( \gamma \) the shear deformation fixed and equal to zero. This is a consequence of the fluctuations being dependent on the ensemble used (see Section 2.4). As the Lebowitz-Percus-Verlet transformation [38] relates the fluctuations in different (conjugated) ensembles, the elastic moduli can be computed using other ensembles than the one chosen here. This issue has been illustrated in some recent works [16, 39].

It is also interesting to point out that the stress fluctuation equations for the elastic moduli, both for the liquid and solid states, can be deduced from general thermostatisical considerations, without introducing a local displacement field, as in ref.[1]. This has been shown in detail in [40] for an isotropic system, where an affine canonical transformation of the positions and the momenta of the particles was made, and the shear modulus obtained via the second derivative of the free energy with respect to the shear deformation, yielding the same equation as eq.(2.124).

### 2.6 Some technical issues

#### 2.6.1 Periodic boundary conditions and minimum images method

In a molecular simulation, the number of particles is always limited. If we only take one simulation box, there will be important surface effects. In order to calculate macroscopic bulk properties of the model system studied, periodic boundary conditions are to be applied [30]. In this way, a small number of particles is extended to an infinite system. This can remove surface effects. As the Fig. 2.1 shows, the shaded box
represents the original simulation box, while the surrounding boxes are exact copies in every detail. Whenever an atom leaves the simulation cell, it is replaced by another with exactly the same velocity, entering from the opposite cell face so that the number of atoms in the cell is conserved.

Fig. 2.1 A two-dimensional simulation box (shaded) containing 5 particles with its nearest periodic images.

In simulations with relatively short range interactions, we only need the nearest neighbours around the simulation box. We calculate the interactions between a particle \( i \) in the box and all the other particles \( j \) within the range of the potential, by applying the minimum image convention [30]. For a given range of the potential, with a cut-off distance, the simulation box side must thus be larger than twice the cut-off distance.

### 2.6.2 Truncation of the pair potential and related corrections

In simulations, it is necessary to truncate the pair potential \( u(r) \) at some finite distance \( r = r_c \) (the cut-off distance), such that \( u(r) = 0 \) for \( r > r_c \). If there is discontinuity of \( u(r) \) at the cut-off distance, then there must be an “impulsive correction” to the virial expression of the pressure [41]. This can be avoided by shifting the pair potential, imposing \( u_{sh}(r) = u(r) - u(r_c) \) for \( r \leq r_c \) and \( u_{sh}(r) = 0 \) for \( r > r_c \). This
shift will not alter the forces, so will not affect the particles trajectories. In this thesis, we use indeed shifted pair potentials. However, this shift will not generally prevent the first derivative of the potential, \( u'(r) \), to be discontinuous at the cut-off. As the Born terms of the elastic constants involve second derivatives of the pair potential, it is necessary to make another “impulsive correction” to these calculations. The issue has been discussed in [42]. Relevant expressions will be given in Chapter 3.

To conclude this chapter, we point out that we have laid out here the theoretical foundation for our applications in the next chapter (Ch.3). They are all concerned with static properties. The time-dependent properties are all gathered in Chapter 4.
Chapter 3

Static elastic properties

In this chapter, we gather static properties computed using the formalism of the previous chapter. We have carried out three investigations. The first is the computation of the bulk moduli of hard sphere crystals by volume fluctuations. The second is a study of a two-dimensional glass-former: the Kob-Andersen 2d model. In the third part we investigate stress fluctuations in model crystals, one is a simple LJ fcc crystal, the other is an AB13 superlattice, formed by a binary mixture of repulsive particles, in the proportion 1:13 of large and small spheres.

3.1 Bulk moduli of hard sphere crystals by volume fluctuations

The bulk modulus is an important property of materials. From its definition, \( K = -V \frac{\partial P}{\partial V} \bigg|_T \), it is possible to compute it by performing a finite (but small) pressure change \( \Delta P \), and measure the change in volume \( \Delta V \), or vice versa. However, this procedure can imply large numerical errors, especially in the regime where the pressure is high, and where it is highly non-linear with respect to the volume change. Within constant pressure simulations, it is possible to evaluate \( K \) from the volume fluctuations during the simulation [30], via the relation \( K = k_B T \frac{\langle V \rangle}{\langle \delta V^2 \rangle} \), where \( k_B \) is the Boltzmann constant, \( \langle V \rangle \) is the average volume at the given thermodynamic state point (NPT) and \( \langle \delta V^2 \rangle = \langle (V - \langle V \rangle)^2 \rangle \) is the variance of the volume (Notice that \( \langle \ldots \rangle \) indicates the thermostatistical averages). This idea is shown to give accurate estimates of \( K \), when compared with other more elaborate stress-fluctuation approaches [35]. But the cases
studied there involve situations with quite weak pressure. Here we apply this method to a hard sphere solid, both monodisperse and polydisperse, up to high pressure, to show first its feasibility and second its predictive power in general physically relevant situations.

The hard sphere crystal phase has been well studied since long time [43]. The analytical fit of the equation of state (EOS) \( \beta P(\rho) \), \( (\beta = 1/k_B T) \), proposed by Young and Alder [44] has proven to accurately cover the whole solid region, and is robust enough when compared to more recent simulations and EOS based on much larger systems [45]. Once the EOS is known, the bulk modulus can be calculated by direct derivative of \( P \) with respect to the density \( \rho \). There exists another route to compute the elastic constants, via the stress-fluctuation formalism [1]. This formalism has been successfully implemented to model glass forming colloidal systems [39]. The special case of hard spheres has been treated by Farago and Kantor [46] for the computation of its elastic constants within the stress-fluctuation formalism. However, the implementation of the method is quite elaborate. Furthermore, the formalism is limited to classical pairwise potentials. This narrows its potential use for some real materials. If only the bulk modulus is needed, then other more direct and simpler simulations can be preferred. In the present work, we show that the widely used constant pressure (\( NPT \)) simulations can indeed produce correctly the bulk modulus \( K \). The method is mostly interesting in cases where an accurate fit of the EOS does not exist, and where the pressure varies quite non-linearly with the density, rendering a finite difference method inaccurate.

### 3.1.1 Monodisperse hard sphere solid

The case of monodisperse hard sphere solid is well known. It will be used here to check the relevance of the method. We perform constant pressure Monte Carlo simulations [30, 41] of hard spheres solid, consisting of an fcc crystal. The simulation box contains \( N = 864 \) hard particles of diameter \( \sigma \). The reduced pressure \( P^* = \beta P\sigma^3 \) varies from about 12 to 50. As \( P \) is proportional to \( T \) (at given \( \rho \)) for hard spheres, the physical controlling parameter is indeed \( \beta P \), instead of \( P \) and \( T \) separately. Although the number of particles is not as large as in some recent simulations [45], it is significant
enough for the purpose of the present work (cf. analysis in [41]). Starting from an fcc crystal, with an initial volume \(V_0\), the system is equilibrated during \(5 \times 10^5\) MC cycles under the constant pressure \(P_0^*\). Statistics are further gathered during typically \(2 \times 10^6\) MC cycles. Information such as the average volume \(\langle V \rangle\), the volume variance \(\langle \delta V^2 \rangle\), and the volume distribution (V-histogram) are the main results we obtain. This allows us to compare our simulation results with the EOS of ref. [44]. For the sake of completeness, we display this EOS in the following form.

\[
\frac{\beta P}{\rho} = \sum_{n=-1}^{3} a_n y^n, \tag{3.1}
\]

where \(y = \rho_{cp}/\rho - 1\) with the close-packing density \(\rho_{cp} = \sqrt{2}\sigma^{-3}\) (fcc structure), and \(a_{-1} = 3, a_0 = 2.566, a_1 = 0.55, a_2 = -1.19\) and \(a_3 = 5.95\). The bulk modulus \(K\) is thus given by the equation

\[
\frac{\beta K}{\rho} = \sum_{n=-2}^{3} b_n y^n. \tag{3.2}
\]

The coefficients \(b_n\) are given in Table 3.1.1. Following our simulation, the first direct result is the average density \(\rho = N/\langle V \rangle\). In Fig. 3.1, we plot \(\rho^* = \rho\sigma^3\) vs \(P^*\) and compare our results with eq.(3.1). We see that indeed, the agreement is excellent. The (reduced) bulk modulus from simulations, \(K^* = \beta K\sigma^3\), is plotted vs \(P^*\) in Fig. 3.2. The simulation results are compared with eq.(3.2). Again, excellent agreement is observed. However, unlike the average density \(\rho\), the bulk modulus \(K\)-results bear some non-negligible measurement uncertainty, which is represented by error-bars in Fig. 3.2. They are estimated by standard deviations of \(K\) from several independent runs. In Fig. 3.3, the volume histogram is shown for a relatively high pressure \(P^* = 35.5\). Indeed, the curve follows a Gaussian distribution, as it should, concerning equilibrium fluctuations [47]. We stress that rather long runs are necessary in order to have Gaussian-like volume histogram, meaning a correct sampling of the volume fluctuations.

### 3.1.2 Polydisperse system

Having shown the feasibility of the method, we try to make some predictions with it. To do this, we take a polydisperse hard sphere system, containing three components,
Fig. 3.1 The reduced average density $\rho^*$ vs the reduced pressure $P^*$ for monodisperse and slightly polydisperse solids. In the monodisperse system, $\rho^* = \rho \sigma^3$ and $P^* = \beta P \sigma^3$. In the polydisperse case, $\rho^* = \rho \bar{\sigma}^3$ and $P^* = \beta P \bar{\sigma}^3$. The continuous line is the analytical EOS of the monodisperse system given by eq.(3.1). The dots represent NPT simulation results of the monodisperse system. The dashed line is the analytical EOS of the polydisperse system given by eq.(3.3), for $P^* \leq 20$. The dotted line is its extrapolation for $P^* \geq 20$. The triangles represent NPT simulation results of the polydisperse system.
Fig. 3.2 The reduced bulk modulus $K^*$ vs the reduced pressure $P^*$ for monodisperse and slightly polydisperse solids. In the monodisperse system, $K^* = \beta K \sigma^3$ and $P^* = \beta P \sigma^3$. In the polydisperse case, $K^* = \beta K \bar{\sigma}^3$ and $P^* = \beta P \bar{\sigma}^3$. The continuous line is given by eq. (3.2). The circles (with error-bars) represent NPT simulation results of the monodisperse system. The dashed line is the analytical expression for the polydisperse system given by eq. (3.3), in the range $12 \leq P^* \leq 20$. The dotted line is its extrapolation for $P^* \geq 20$. The triangles represent NPT simulation results of the polydisperse system (error-bars, similar to the monodisperse case, are not shown).
Table 3.1.1 Coefficients entering in Eqs.(3.1) and (3.2). \( \{a_n\} \) are taken from the work of Young et al. [44], \( \{b_n\} \) are computed using \( K = \rho \frac{\partial P}{\partial \rho} \). \((b_{-2} = -a_{-1}, \ b_3 = -2a_3, \) and \( b_n = (1 - n)a_n - (1 + n)a_{n+1} \) for \( n = -1 \) to 2).

<table>
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<td></td>
</tr>
<tr>
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<td>2.38</td>
<td>-16.66</td>
<td>-11.9</td>
</tr>
</tbody>
</table>

with the composition 1/3 for each of them, and diameters \( \sigma_1 = (1 - \epsilon)\sigma_0 \), \( \sigma_2 = \sigma_0 \) and \( \sigma_3 = (1 + \epsilon)\sigma_0 \). The mean diameter (the first moment of the size-distribution) \( \bar{\sigma} = \sigma_0 \), the polydispersity index \( I = \frac{\sigma^2}{\bar{\sigma}^2} - 1 = (2/3)\epsilon^2 \) (\( \sigma^2 \) is the second moment of the size-distribution). This system has been studied previously by one of us, in the framework of the crystallisation [48]. Here we do not consider the fluid phase, because for this phase, the well-known EOS by Mansoori et al. provides a quite accurate description of the polydisperse fluid, at least up to the crystallisation densities. The crystallisation is out of our scope too, because of the fractionation phenomenon accompanying the freezing of polydisperse systems [48]. We thus limit ourselves to the crystal phase, in the same range of the reduced pressure as for the monodisperse system, i.e. \( 12 \leq P^* \leq 50 \) (\( P^* = \beta P \bar{\sigma}^3 \)). For low to moderate (reduced) pressure, i.e. \( 1.0 \leq \rho \bar{\sigma}^3 \leq 1.2 \), an EOS has been proposed by Bartlett [49], based on the hypothesis that the thermodynamics of a polydisperse systems can be mapped to that of a binary mixture (through the identification of the first three moments of the size distribution), and taking advantage of the extensive simulations on binary hard sphere solid make by Kranendonk and Frenkel [50, 51]. This EOS (eq.(34) of ref. [49]) leads naturally to a approximate analytical expression for the bulk modulus \( K \), which we shall test with our simulation results. The EOS of ref. [49] contains 20 double-precision fitting coefficients. In the case of a symmetric size-distribution, the number of coefficients is reduced to 9. This number is still large, compared to the monodisperse case, showing the difficulty of fitting the EOS of polydisperse high density systems. In the following, we display the

40
EOS of ref. [49], in the case of symmetric size-distribution (relevant to our system).

\[
\beta P_p \bar{\sigma}^3 = \beta P_m \bar{\sigma}^3 + \sum_{i,j} C_{ijk} \rho \bar{\sigma}^{3i} (10^4 I)^j
\]  

(3.3)

where \( P_p \) stands for the pressure of the polydisperse system, \( P_m \) that of the monodisperse system (given by eq.(3.1)) as a function of the dimensionless density \( \rho \bar{\sigma}^3 \), the coefficients \( C_{ijk} \) are displayed in Table 3 of ref. [49], and \( I \) is the above-defined polydispersity index. From eq.(3.3), the bulk modulus \( K \) is readily expressed to be

\[
\beta K_p \bar{\sigma}^3 = \beta K_m \bar{\sigma}^3 + \sum_{i,j} i C_{ijk} \rho \bar{\sigma}^{3i} (10^4 I)^j
\]  

(3.4)

where the \( K_p \) and \( K_m \) stand respectively for the bulk modulus of the polydisperse and monodisperse systems. \( K_m \) is given by eq.(3.2). Our simulated system consists always of \( N = 864 \) particles, with 288 particles for each species. In this work, a weak polydispersity is taken, i.e. \( \epsilon = 0.02 \). This will allow us to investigate how \( \rho \) and \( K \) are sensitive to the polydispersity, at given \( P^* \). In Fig. 3.1, we plot \( \rho^* = \rho \bar{\sigma}^3 \) vs \( P^* \) for the polydisperse system. Two points of observation can be made. First, \( \rho \) is indeed sensitive to the polydispersity. At a given pressure, the density is sensibly lower in a polydisperse system. This is consistent with the commonly known fact that the crystal is destabilized by polydispersity (see e.g. ref [50]). The second observation is that the EOS of ref. [49] coincide with our simulation results in the range of its initial domain of fit. And not surprisingly, it deviates from simulation results outside this domain. So extrapolation of this EOS is not correct. In Fig. 3.2, we have plotted \( K^* = \beta K \bar{\sigma}^3 \) vs \( P^* \) for the polydisperse system. Here again, we see that the analytical expression eq.(3.4) cannot be extrapolated beyond \( P^* \sigma^3 \approx 20 \). On the other hand, at given \( P^* \), the value of \( K^* \) do not differ much with the monodisperse case. This result can be interpreted by two opposite effects of the polydispersity. Actually, in a (slightly) polydisperse solid, at given \( P^* \), \( \langle V \rangle \) is larger, but \( \langle \delta V^2 \rangle \) too. These two effects somehow compensate each other in \( K \). In Fig. 3.3, the volume histogram is shown at a rather high (reduced) pressure \( P^* = 35.5 \). Compared to the monodisperse case, the distribution is shifted to higher volumes, and is slightly broadened. Again, a fit with Gaussian distribution represents accurately the histogram.
Fig. 3.3 Volume-histogram at the reduced pressure $P^* = 35.5$ for monodisperse and slightly polydisperse solids. The dots represent $NPT$ simulation results of the monodisperse system. The triangles represent those of the polydisperse system. The continuous and dashed lines represent fits by Gaussian distribution for respectively the monodisperse and polydisperse cases. $V^* = V/\sigma^3$ for the monodisperse case, and $V^* = V/\bar{\sigma}^3$ for the polydisperse case. The histograms are scaled such that their maximum value is one.
3.1.3 Topical summary

In summary, we tested the feasibility of directly computing the bulk modulus from \(NPT\) simulations, by taking the hard sphere solids. The method can cover a very large range of the solid phase, including high pressure solid, where finite difference approach is not appropriate. In the polydisperse case, we show first that the EOS is sensitive to the polydispersity, and secondly, the EOS of Bartlett cannot be extrapolated beyond its initial fitting range (of low to moderate pressures). By computing \(K\) up to \(P^* = 50\), we demonstrate that these simulations can indeed provide correct \(K\) values for high pressure polydisperse solids. Finally, we point out two opposite effects of the polydispersity, rendering \(K\) much less sensitive to the polydispersity than the average density does (at a given pressure). This method can find many applications for various systems and situations, for example in the case of structural phase transition in metallic systems, or in the case of self-organisation in complex liquids.

3.2 A glass former in two dimensions: 80-20 Kob-Andersen model

3.2.1 Background

The classical 80-20 Kob-Andersen model (KA) [19], a binary Lennard-Jones (LJ) bead mixture with a fraction \(f = 0.8\) of large spheres, is an important numerical reference model for the understanding of the glass transition in three dimensions [9, 12, 52–54]. Its recently proposed two-dimensional (2d) version, called below “KA2d model”, has been much less investigated [55], despite the high experimental relevance of 2d glass-forming systems [28, 56–60]. This may be due to the disappointing finding reported in [55] that the standard KA fraction of large spheres did not show a glass transition, but rather a crystal-like low-temperature phase. Various different fractions \(f\) have thus subsequently been studied [55, 61–69] with the notable exception of [62] where the classical 80-20 KA model has also briefly been considered in two dimensions. In the present work we readdress some of the results reported in [55]. Here are general goals of present work. One central point we want to make is merely that most computational studies [55, 61–65, 67–69] compare configurations prepared and sampled at an imposed
constant volume \( V \) and this in addition corresponding to an unrealistically large number density \( \rho \approx 1.2 \) (in units of the large sphere’s diameter) [55]. Obviously, from the experimental and application point of view one should rather control the pressure \( P \) (or more generally the stress tensor) while the systems are quenched into the amorphous state. Surprisingly, there exist at present only a few computational studies [8, 66] where the 2d glass transition is investigated allowing the volume to fluctuate at an imposed constant pressure. Following our recent study of the three-dimensional (3d) KA model (KA3d) [70], the aim of the current work is thus to characterize the standard KA2d model at a moderate pressure corresponding to a much smaller density \( \rho \approx 1.0 \) in the low-temperature limit. We show that under these conditions the KA2d model is in fact a reasonably good glass-former, just as its 3d counterpart, and no indications of a crystallization or other forms of long-range orientational correlations [69] have been found. As a companion system we investigate in parallel a monodisperse Lennard-Jones (mdLJ) system under the same external constraints. As one expects [36], this model forms a triangular lattice below a freezing temperature \( T_f \). We compare thus various properties of the KA2d model glass with this reference. There exists a cusp-singularity of shear modulus. The present study focuses on simple static and quasi-static thermodynamic properties such as the number density \( \rho(T) \), the compression modulus \( K(T) \) or the shear modulus \( G(T) \) as a function of the temperature \( T \) on both sides of the glass transition temperature \( T_g \). The equilibrium shear modulus \( G \) is obviously an important order parameter characterizing in general the transition from the liquid/sol \((G = 0)\) to the solid/gel state \((G > 0)\) where the particle permutation symmetry of the liquid state is lost for the time window probed [71, 72]. (Please note that all reported shear moduli are quasi-static or transient in this sense.) As in other related numerical studies [2, 8, 16, 40, 42, 70, 73] we shall determine the only two relevant elastic moduli \( K(T) \) and \( G(T) \) using the stress-fluctuation formalism [1, 4, 9, 30, 41]. In contradiction to the additive jump discontinuity predicted by the mode-coupling theory [28, 53, 74, 75], we show that the shear modulus \( G(T) \) of the KA2d model reveals a continuous cusp-singularity

\[
G(T) \approx G_g (1 - T/T_g)^{\alpha} \quad \text{for} \quad T/T_g < 1 \tag{3.5}
\]
with $\alpha \approx 0.6$. This result is in qualitative agreement with related numerical [2, 39] and theoretical [76] work, albeit in conflict to some recent experimental work [28, 59].

We begin by presenting in subsection 3.2.2 the numerical model systems considered and summarize the relevant stress-fluctuation relations for the determination of $K(T)$ and $G(T)$. We describe then in subsection 3.2.3 our simulation results and summarize finally this work in subsection 3.2.4.

### 3.2.2 Algorithmic Details

**Model Hamiltonians.** For comparison we have studied two soft bead models. As a reference we compute systems of monodisperse Lennard-Jones beads (mdLJ) [30], i.e. we take advantage of a generic model for “simple liquids” [52] with a perfectly known 2d phase diagram [36]. Our system consists of $N = 1250$ particles interacting through a shifted and truncated LJ potential. The central model of the present study is the 2d version (KA2d) [55] of the standard 3d KA model [19]. It consists of a binary mixture of LJ particles. We have sampled these model Hamiltonians by means of molecular dynamics (MD) simulations [30, 41] using the LAMMPS code [29] and taking advantage of the standard Nosé–Hoover thermostat and barostat provided by this algorithm. The temperature coupling constant is set to 10 and the isobaric coupling constant to 25 (with both values in simulation units). A rectangular box with $L_y = \sqrt{3}L_x$ was used in order to be compatible with the triangular crystal phase of the mdLJ model. For both models the average normal pressure $P$ is kept at a constant $P = 2$ for all temperatures considered as in [70]. For the mdLJ model we have carried out both cooling and heating cycles. For the KA2d model the systems have been quenched with a constant cooling rate of $10^{-5}$ starting from the liquid limit at $T = 1$. This corresponds to the main rate discussed in [55]. We remind that areas of hexagonally crystallized A particles in a matrix of amorphous AB material were observed for exactly this rate and the standard fraction $f = 0.8$ of large spheres. Table 3.2.1 summarizes some properties of the KA2d model discussed in the next section. We use stress-fluctuation relations for elastic moduli. The compression modulus $K(T)$ and the shear modulus $G(T)$ indicated in table 3.2.1 have been obtained using the relevant stress-fluctuation relations for simulations at constant volume $V$ and constant shear strain.
\( \gamma = 0 \) [1, 4, 30, 41, 70]. While the systems are quenched imposing a constant mean pressure, i.e. the volume fluctuates while we cool, the elastic moduli are subsequently determined for each T in the NVT-ensemble by switching off the barostat after some tempering [8]. The compression modulus is given by the Rowlinson stress fluctuation formula [35, 41]

\[
K = \eta_{A,ex} - \eta_{F,ex} + P_{id} \tag{3.6}
\]

The first “affine” contribution \( \eta_{A,ex} \) is a sum of moments of first and second derivatives of the pair potentials used. Characterizing the average energy change under an affine dilatation strain it is related to the well-known “hypervirial” [30]. See [35, 70] for details.

The second contribution \( \eta_{F,ex} \) stands for the reduced fluctuation \( \eta_{F,ex} \equiv \beta V \langle \delta \hat{P}_{ex}^2 \rangle \) of the instantaneous excess pressure \( \hat{P}_{ex} \) (with \( \beta \) being the inverse temperature) and the third contribution \( P_{id} \) to the mean ideal pressure. As in related studies [2, 8, 16, 24, 40, 42, 70] the shear modulus \( G \) may be obtained most readily using

\[
G = \mu_{A,ex} - \mu_{F,ex} = \mu_A - \mu_F \tag{3.7}
\]

with \( \mu_{A,ex} \) being the excess contribution to the affine shear elasticity \( \mu_A = \mu_{A,id} + \mu_{A,ex} \), a simple average of moments of first and second derivatives of the pair potential characterizing the mean energy under an affine pure shear strain. The second contribution \( \mu_{F,ex} \equiv \beta V \langle \delta \hat{\tau}_{ex}^2 \rangle \) stands for the excess contribution to the total shear stress fluctuation \( \mu_F = \mu_{F,id} + \mu_{F,ex} \) with \( \hat{\tau}_{ex} \) being the excess contribution to the instantaneous shear stress. Since for a shear strain at constant volume the ideal free energy contribution does not change, the explicit kinetic energy contributions must be irrelevant for \( G \). (An ideal gas can not elastically support a finite shear stress.) As one thus expects, the kinetic contributions \( \mu_{A,id} = \mu_{F,id} = P_{id} \) cancel and can be dropped when \( G \) is determined using the first equation of eq.(3.7). We note finally that since second derivatives are relevant for \( \eta_{A,ex} \) and \( \mu_{A,ex} \), impulsive corrections must be taken into account for the truncated and shifted potentials considered in the present work as stressed in [42]. Otherwise the shear modulus \( G \), determined using eq.(3.7), does not vanish in the liquid limit as it must.

### 3.2.3 Numerical results

**Specific volume.** As explained above, we impose a normal pressure \( P = 2 \) for all
Table 3.2.1 Several properties of the KA2d model at pressure $P = 2$ as a function of temperature $T$: number density $\rho = N/V = 1/v$, linear box size $\sqrt{3}L_x = L_y$, Abraham parameter $R = g_{\text{min}}/g_{\text{max}}$ [77], ideal pressure contribution $P_{\text{id}} = T\rho$ to the imposed normal pressure $P = P_{\text{id}} + P_{\text{ex}}$, hypervirial contribution $\eta_{A,\text{ex}}$ to the compression modulus, compression modulus $K$ obtained using the Rowlinson formula eq.(3.6) [30, 35, 70], excess contribution $\mu_{A,\text{ex}}$ to the affine shear elasticity $\mu_A = \mu_{A,\text{id}} + \mu_{A,\text{ex}}$ and shear modulus $G$ obtained using the stress-fluctuation formula eq.(3.7). The last five columns have the dimension energy per volume. The error bars for $K$ and $G$ are of order one in these units, all other properties are known to higher precision. Boltzmann’s constant is set to unity and all properties are given in LJ units [30].

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<td>0.82</td>
<td>42.7</td>
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</table>

systems considered. If the systems are cooled down, the number density $\rho(T)$ must thus increase (Table 3.2.1) and the specific volume $v(T) = 1/\rho(T)$ per particle must decrease. This is shown in Fig. 3.4 for both models. A jump singularity is observed for $v(T)$ at the freezing temperature $T_f \approx 0.62$ of the mdLJ model. This jump is of course a consequence of the first order phase transition from the liquid phase to the crystalline state [1, 33, 36]. At variance with this, the KA2d model changes more gradually with linear slopes fitting reasonably both the low- and the high-T limits. The observed (more or less) sudden change of the tangent slopes at the glass transition may be used to operationally define $T_g$ [12] by matching the indicated two lines as done in
Fig. 3.4 Rescaled specific volume $v(T)$ vs. temperature $T$ for both models considered. The freezing temperature $T_f \approx 0.62$ for the mdLJ model and the glass transition temperature $T_g \approx 0.35$ for the KA2d model are indicated by vertical lines.

Various recent numerical studies [8, 55, 70]. Using this calorimetric criterion we obtain $T_g \approx 0.35$ for the KA2d model. This is slightly smaller than the corresponding value $T_g \approx 0.41$ for the KA3d model at $P = 1$ [70]. We remind that for the poly-disperse purely repulsive LJ system (pdLJ) considered in [5, 42, 70, 78] the same criterion yields a glass transition temperature of only $T_g \approx 0.26$ at the same pressure $P = 2$. In qualitative agreement with [61] this suggests that, having a much larger $T_g$, the KA2d model should be a more promising numerical model for investigations of the glass transition in two dimensions.

**Radial distribution functions.** One of the conclusions of [55] is that the KA2d model with a fraction $f = 0.8$ of large spheres should crystallize instead of forming an amorphous glass at low temperatures. In order to clarify this issue for our constant pressure systems, we compare in the main panel of Fig. 3.5 the radial pair-distribution function (RDF) of the mdLJ crystal at $T = 0.3$ (spheres) with the (total) RDF $g(r)$ of the KA2d system for $T = 0.1$ (solid line). Whereas the former system becomes clearly
crystal-like with long-range spatial correlations (at least if boundary conditions with $L_y = \sqrt{3}L_x$ are used), we find for the KA2d model that $g(r) \approx 1$ for $r > 4$ as is typical for an amorphous substance. The amorphous, non-crystalline behaviour is also confirmed using the corresponding structure factor $S(q)$ in reciprocal space (not shown) and more readily by inspection of snapshots such as the one for $T = 0.2$ shown in the inset of Fig. 3.5. We use R-parameter criterion. An additional empirical criterion for the liquid-to-solid/glass transition has been proposed by Abraham [77]. It uses the temperature dependence of the so-called R-parameter defined by $R \equiv g_{\text{min}}/g_{\text{max}}$ with $g_{\text{max}}$ being the main peak of the radial pair-distribution function (RDF) and $g_{\text{min}}$ its subsequent minimum. As shown in Fig. 3.6, R(T) shows for the mdLJ model a jump discontinuity at $T_f \approx 0.62$ consistent with Fig. 3.4. Although the criterion was suggested originally only for a one-component system, we extend it here to our binary mixture in two ways by using either the total RDF $g(r)$ sampled over all beads or the RDF $g_{11}(r)$ considering only the large spheres. The R-parameter obtained from the latter distribution is shown by the diamonds. The glass transition temperature may be obtained by matching the low- and the high-temperature linear slopes indicated in the figure. We get $T_g \approx 0.35$ as above from the specific volume $v(T)$.

**Compression modulus.** We turn now to the two elastic moduli, the compression modulus $K(T)$ and the shear modulus $G(T)$, which characterize completely the linear elastic response of both models even in the triangular phase of the mdLJ model. The temperature dependence of the compression modulus $K$ (circles) and its contributions $\eta_{A,ex}$ (squares) and $\eta_{F,ex}$ (triangles) is presented in Fig. 3.7. The affine hypervirial contribution $\eta_{A,ex}(T)$ is seen in both cases to increase more or less linearly with decreasing temperature. As seen in panel (b) for the KA2d model, one may determine again a transition temperature from the intercept of the low- and the high-T tangent slopes. This criterion suggests a glass transition temperature $T_g \approx 0.32$ which is consistent, albeit slightly smaller than the one obtained above. The excess normal pressure fluctuation contribution $\eta_{F,ex}(T)$ shows a striking peak at the freezing temperature $T_f \approx 0.62$ of the mdLJ model and vanishes then rigorously for $T \to 0$. Since in this limit the ideal pressure contribution $P_{id}$ vanishes also, the compression modulus $K(T = 0)$ is completely determined from the affine contribution $\eta_{A,ex}$. This is expected for sim-
Fig. 3.5 Main panel: Radial pair-distribution function $g(r)$ vs. distance $r$ for two low-temperature states of both models. While long-range correlations are visible for the mdLJ model (small circles), no long-range order is seen for the KA2d model where $g(r) \to 1$. Inset: Snapshot for $T = 0.2 \ll T_g$ confirming the amorphous structure of the KA2d model. Open/filled circles correspond to the large/small beads.
Fig. 3.6 R-parameter of both models vs. temperature T. A jump discontinuity at $T_f \approx 0.62$ is seen for the mdLJ model. The linear low- and high-temperature linear slopes of the KA2d model match again at $T_g \approx 0.35$. 

$P=2$

$R=0.03+0.12T$

$T_g=0.35$

$T_f=0.62$

$T_g=0.05+0.35T$
ple lattice models with one atom per unit cell [73]: if the zero-temperature triangular lattice is strained in an affine manner by changing the simulation box, all beads remain mechanically stable and there are no non-affine displacements relaxing the strain energy. As seen in panel (b), this is different for the KA2D model where $\eta_{F,ex}(T)$ decreases below $T_g \approx 0.32$ but does not vanish at $T = 0$ as indicated by the horizontal arrow. As explained in detail, e.g., in section 4 of [40], the forces on the particles of an affinely strained amorphous body do normally not vanish which leads to non-affine displacements lowering the strain energy and, hence, the modulus. It is for this reason that the affine contribution $\eta_{A,ex}(T = 0) \approx 88$ only sets an upper bound to the modulus $K(T = 0) \approx 84$ with $\eta_{F,ex}(T = 0) \approx 4$ measuring the energy reduction due to the non-affine displacements. The compression modulus $K(T)$ of the KA2d model thus increases strongly below $T_g \approx 0.32$ but does not reach $\eta_{A,ex}(T)$ as is the case for the mdLJ model at $T = 0$.

**Shear modulus as a function of temperature.** Our numerical results for $\mu_{A,ex}(T)$, $\mu_{F,ex}(T)$ and $G(T)$ are presented in panel (a) of Fig. 3.8 for the mdLJ model and in panel (b) for the KA2d model. Note that the simple average $\mu_{A,ex}(T)$ is continuous in both cases increasing monotonously with decreasing temperature, as may be expected due to the increasing density, with similar (albeit not identical) numerical values for all temperatures. (The observed change of the tangent slope for the KA2d model is too weak to allow a precise determination of $T_g$.) For both models the stress fluctuation contribution $\mu_{F,ex}(T)$ to the shear modulus $G$ is identical in the liquid high-T limit to the affine shear elasticity $\mu_{A,ex}$. This is expected from eq.(3.7) and the fact that the shear modulus of a liquid must vanish. $\mu_{F,ex}(T)$ shows a maximum at, respectively, $T_f \approx 0.62$ for the liquid-solid and $T_g \approx 0.32$ for the liquid-glass transition. While $\mu_{F,ex}(T)$ vanishes rigorously in the low-T limit for the mdLJ model due to the simple triangular lattice adopted [73], $\mu_{F,ex}(T)$ is seen to level off for the KA2d model (horizontal arrow). In agreement with the discussion of the compression modulus $K$ above, this confirms that an affine shear does neither lead for an amorphous body to a mechanically stable configuration and a finite energy per volume $\mu_{F,ex}(T = 0) \approx \mu_{A,ex}(T = 0)/2$ of the strained ground state system can be relaxed by non-affine displacements as discussed elsewhere [5, 16, 40, 42, 70, 73, 78]. As one
Fig. 3.7 Compression modulus $K(T)$ and its contributions $\eta_{A,ex}(T)$ and $\eta_{F,ex}(T)$ according to the Rowlinson stress-fluctuation formal eq.(3.6) : (a) Data for the mdLJ model. $K(T)$ vanishes at the phase transition at $T_f \approx 0.62$. The stress fluctuation contribution $\eta_{F,ex}(T)$ vanishes for $T \to 0$. (b) Data for the KA2d model showing a similar, albeit slightly smaller transition temperature $T_g \approx 0.32$ as above. $\eta_{F,ex}(T)$ does not vanish for $T \to 0$. 
expects for the mdLJ model, a clear jump singularity at $T_f \approx 0.62$ is observed in panel (a) for the stress fluctuation contribution $\mu_{F,ex}$ and (consequently) for the shear modulus $G = \mu_{A,ex} - \mu_{F,ex}$. At striking variance with this a continuous behaviour is observed for $\mu_{F,ex}(T)$ and $G(T)$ at the glass transition of the KA2d model. Confirming qualitatively the cusp-singularity found in [70], a power law with $G = G_g(1 - T/T_g)^\alpha$ and $G_g \approx 16.5$, $T_g \approx 0.32$ and $\alpha \approx 0.6$ seems to fit our data. This is better seen from the zoom on the glass transition region given in the inset of Fig. 3.8 (b). We remind that a slightly lower exponent $\alpha \approx 1/2$ was fitted for the KA3d model in our previous work [70] as may be seen from the rescaled data (diamonds) included in the inset. Whether the weak difference of the exponents reflects a fundamental effect due to the different spatial dimensions, as suggested by the recent work of Flenner and Szamel [69], or whether it should merely be attributed to the small system sizes used, can currently not be answered. Larger systems and, more importantly, longer trajectories are warranted to clarify this question.

**Sampling time dependence of stress fluctuations.** As already emphasized elsewhere [8, 16, 70], the convergence with sampling time of the stress fluctuation relations for the elastic moduli eqs. (3.6) and (3.7) may be slow even for permanent elastic networks and one thus has to check whether sufficiently large trajectories have been used to determine reliable long-time estimates. The reason for this is that, while the simple means $\eta_{A,ex}$ and $\mu_{A,ex}$ converge essentially immediately, the stress fluctuation contributions $\eta_{F,ex}$ and $\mu_{F,ex}$ become sampling-time dependent properties $\bar{\eta}_{F,ex}(\Delta t)$ and $\bar{\mu}_{F,ex}(\Delta t)$ if sampled over trajectories of finite lengths $\Delta t$. Please note that this has apriori nothing to do with an insufficient equilibration or ageing of the systems, but stems from the fact that the stress-fluctuations simply need time to explore the available phase space. Our results for this topic will be shown in the next chapter, with other time-dependent properties. We note here briefly that indeed, a strong dependence has been observed, especially around the glass transition.

### 3.2.4 Topical conclusions

To summarize, we have investigated by means of molecular dynamics simulations the glass transition of the two-dimensional version of the Kob-Andersen model with a
Fig. 3.8 Affine shear elasticity $\mu_{A,ex}$ (squares), shear stress fluctuation $\mu_{F,ex}$ (triangles) and shear modulus $G = \mu_{A,ex} - \mu_{F,ex}$ (circles) vs. temperature $T$: (a) For mdLJ systems $G(T)$ reveals a striking jump discontinuity at $T_f \approx 0.62$. (b) The continuous increase of $G(T)$ for KA2d systems below $T_g \approx 0.32$ shows a cusp-singularity with $\alpha \approx 0.6$. $\mu_{F,ex}$ (triangles) does not vanish for $T \to 0$. Inset: Zoom on the glass transition region plotting $G(T) = G_g$ vs. $x = T/T_g$ for the KA2d model ($T_g = 0.32, G_g = 16.5$) and the KA3d model ($T_g = 0.41, G_g = 24.5$). The lines indicate equation eq.(3.5) with $\alpha = 0.6$ for the KA2d model and $\alpha = 0.5$ for the KA3d model.
fraction $f = 0.8$ of large spheres (as in the standard 3d KA model [19]. In contrast with other recent studies [55, 61–65, 67–69], one moderate mean pressure of order unity has been imposed for all temperatures considered. We have compared our KA2d model with a simple companion model provided by monodisperse LJ beads (mdLJ). Using calorimetry (Fig. 3.4), the radial pair-distribution function (Fig. 3.5), the R-parameter characterization of the radial pair-distribution function (Fig. 3.6), the compression modulus $K$ (Fig. 3.7) and the shear modulus $G$ (Fig. 3.8) it has been demonstrated that this KA2d version is a reasonably good glass-former with a glass transition temperature $T_g \approx 0.3$. Interestingly, as in [70] a slightly smaller glass transition temperature $T_g$ is fitted from the elastic moduli ($T_g \approx 0.32$) as from the specific volume or the R-parameter ($T_g \approx 0.35$). We emphasized that the computation of the shear modulus using the stress-fluctuation relation eq.(3.7) depends strongly on the sampling time $\Delta t$. This makes the determination of the equilibrium shear modulus of the amorphous solid below $T_g$ more delicate than the one of a simple crystal. As one expects, $G(T)$ reveals for the mdLJ model a jump singularity at the freezing temperature $T_f \approx 0.6$. More importantly, we observe for the KA2d model a continuous cusp-singularity below the glass transition temperature (Fig. 3.8 b) confirming thus recent numerical work using similar glass-forming colloidal systems in two and three dimensions [70].

**Discussions.** We note here that all the presented plateau values $G(T)$, especially for the KA2d model [79], correspond strictly speaking to intermediate (albeit perhaps rather large) shoulders of $\mu_{A,ex} - \bar{\mu}_{F,ex}(\Delta t)$. Note that $G(T)$ is only finite below $T_g \approx 0.3$ since on the time scale $\Delta t$ available numerically the particle permutation symmetry of the liquid state is lost [71]. As discussed in [70], if trajectories of increasing lengths are compared, $G(T)$ becomes monotonously smaller, $T_g$ is shifted to smaller values and the glass transition becomes sharper. The phenomenological exponent $\alpha(\Delta t)$ characterizing the cusp-singularity eq.(3.5) should thus decrease somewhat with the sampling time $\Delta t$. It is thus possible that the jump discontinuity ($\alpha \to 0$) predicted by mode coupling theory may be recovered by appropriately extrapolating $\alpha(\Delta t)$ for larger sampling times $\Delta t$.

It is of course necessary to investigate in future work larger systems. Preliminary tests with systems comprising $N = 10000$ particles suggest, however, that all
the reported properties are system size independent, especially no indication for a crystallization has been observed. Even more importantly, as already stated above, future studies should focus on sampling time effects. This should also allow to verify whether the amorphous low-temperature state (Fig. 3.5) remains stable and no additional static long-range correlations (of perhaps Kosterlitz-Thouless-type as suggested in [69]) may become relevant. In view of recent experimental trajectory measurements on 2d glass-forming colloidal systems [28, 58–60] it is also important to analyse the Fourier transform of the displacement fields and other wavevector dependent static and dynamical properties [69].

3.3 Shear stress fluctuations in simple and complex crystals

In this section we calculate the elastic constants of two model crystals, using the stress fluctuation formalism. The two model are the Lennard-Jones system, and a binary mixture of repulsive particles in 1:13 portion, the AB13 system. Our focus will be mainly to point out the importance of the fluctuation terms in crystals, since this term is usually neglected in classical calculations of the elasticity of crystals [7].

3.3.1 Lennard-Jones model system

The Lennard-Jones model is the most popular one in molecular simulations. At low temperatures, it forms a fcc crystal [80]. It is the one used by Squire et al. for the first calculation of the elastic constants in solids by the stress fluctuation formalism [1]. We revisit here the model by considering it as a simple crystal.

Simulation details

In our molecular dynamics simulation, we use NPT and NVT ensembles. The Lennard-Jones potential between a pair of particles at a distance \( r \) is

\[
u_{LJ}(r) = 4\epsilon \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^{6} .
\]

(3.8)
In practice, a shifted version is used:

\[ u_{LJsh}(r) = \begin{cases} 
4\epsilon \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^{6} - u_0, & r < r_c \\
0, & r \geq r_c 
\end{cases} \]  

where \( r_c \) is the cutoff distance of the potential, and \( u_0 = u_{LJ}(r_c) \). All units are reduced by \( \sigma \), \( \epsilon \) and \( m \) (\( m \) being the mass of a particle). We chose \( r_c = 2.5 \) in our simulations. For the heating or cooling or equilibrium processes, NPT ensemble is used while for the statistical calculations NVT ensemble is used. At the beginning, the configuration of our system is obtained at \( T = 0.4 \). Then we have done two types of work including cooling and heating. The processes are as follows: First, cooling or heating the system. We cool or heat the system from the initial configuration (\( T = 0.4 \)) to the next temperature (for example, \( T = 0.3 \) or \( T = 0.5 \)) using NPT ensemble and the cooling or heating rate is \( 10^{-5} \). The MD integration time step is 0.005 in reduced units. Second, we equilibrate the system by \( 2 \times 10^6 \) timesteps using NPT ensemble. In order to keep the pressure at the fixed value (here 1.0), we calculate the average volume during the equilibrium process and change the volume of the last configuration of the equilibrium process to this average volume before running again \( 1 \times 10^6 \) timesteps. The MD integration time step is still 0.005 in reduced units. Third, we collect the statistics by running \( 2 \times 10^6 \) timesteps using NVT ensemble. The MD integration time step is 0.0025 in reduced units.

On the one hand, the cooling process of the configurations as the Table 3.3.1 shows. At every temperature point, the same three steps are repeated. On the other hand, we heat the configuration at \( T = 0.4 \) to the configuration at \( T = 0.5 \) and the heating rate is \( 10^{-5} \), then we reproduce the same three steps as above. The same run is done for other temperatures as the Table 3.3.1 shows. By checking the specific volume (see below), we see that the melting temperature \( T_m \) is located at \( 0.8 < T_m < 0.85 \). Thus more temperature points are examined in this range.

### 3.3.2 Numerical results

**Specific volume.** We first examine the evolution of the specific volume \( v = V/N \) as a function of the temperature. From the graph Fig. 3.9, we can see that there is a
Table 3.3.1 Lennard-Jones system. Temperature points studied by cooling or heating a reference equilibrium system at $T = 0.4$ (crystal). The pressure is $P = 1$ throughout.

<table>
<thead>
<tr>
<th>Process</th>
<th>Temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cooling</td>
<td>0.4 0.3 0.2 0.1 0.05 0.01</td>
</tr>
<tr>
<td>Heating</td>
<td>0.4 0.5 0.6 0.7 0.8 0.81</td>
</tr>
<tr>
<td></td>
<td>0.81 0.82 0.83 0.84 0.85 0.86</td>
</tr>
<tr>
<td></td>
<td>0.86 0.88 0.9 1.0 1.1 1.2</td>
</tr>
</tbody>
</table>

Specific volume for LJ model system

$N = 1372, P = 1.0, T = 0.01-1.2$

Fig. 3.9 The specific volume as a function of the temperature for LJ model system. The pressure is fixed at $P = 1$.

jump along the curve $v(T)$, which indicates the phase transition from crystal to liquid for LJ model system. The melting (freezing) temperature is about $T_m = 0.84$.

**Structural information.** We studied the radial distribution function of the system for several temperatures. From the graph Fig. 3.10, we can see that when $T = 0.1$, the LJ model system shows that it has the regular long distance structure as the peaks
Fig. 3.10 Radial distribution function for LJ model system at different temperatures (T= 0.1, 0.83, 0.84,1.0), from the most structured to the less structured curves. Note that the melting temperature is \( T_m = 0.84 \) (see Fig. 3.9).

indicate. When \( T = 1.0 \), the LJ model system shows that it is in disorder at long range as the radial distribution function at long distances is close to 1. The radial distribution function changes qualitatively from \( T = 0.83 \) to \( T = 0.84 \), which corresponds to the phase transition. The temperature for the phase transition is about 0.84. This confirms the results above based on the specific volume.

**Elastic constants.** For cubic crystals, only three of the \( C_{\alpha\beta\chi\kappa} \) elastic constants are nonzero and independent as follow : \( C_{11}, C_{12}, C_{44} \) where Voigt notation is used. We refer to the work of B. Schnell et al.\[8\] and have obtained the elastic constants and fluctuation terms of the elastic constants for LJ model system and their dependence on temperature at NVT ensemble. The results are as follow. \( C_{11} \) is the average of \( C_{xxxx} \), \( C_{yyyy} \) and \( C_{zzzz} \) and \( C_{12} \) is the average of \( C_{xxyy}, C_{xxxz} \) and \( C_{yyzz} \). \( C_{44} \) is the average of \( C_{xyxy}, C_{xzzz} \) and \( C_{yzyz} \). During the calculation, impulsive correction is imposed on the
The impulsive correction formula is as follows:

\[ C^{B}_{\alpha\beta\chi\kappa} = C^{B,\text{cut}}_{\alpha\beta\chi\kappa} + \Delta C^{B}_{\alpha\beta\chi\kappa}, \]

where \( C^{B,\text{cut}}_{\alpha\beta\chi\kappa} \) are results without impulsive corrections, and \( \Delta C^{B}_{\alpha\beta\chi\kappa} \) are the impulsive corrections. For a cubic crystal, they can be written, in an "isotropic" approximation, as

\[
\begin{align*}
\Delta C^{B}_{44} &= \Delta \mu_B, \\
\Delta C^{B}_{11} &= 3 \Delta \mu_B, \\
\Delta C^{B}_{12} &= \Delta \mu_B
\end{align*}
\]

where

\[
\Delta \mu_B = -\frac{8\pi^{3/2}}{3\Gamma(3/2)} (\rho \sigma^3)^2 \frac{\epsilon}{\sigma^3} f(r_c) g(r_c)
\]

with \( f(r_c) = [1 - (r_0/r_c)^6]/r^3 \) with \( r_0 = 2^{1/6} \) at which the potential \( u_{LJ} \) is minimum. The \( \Gamma(3/2) \) is a gamma function and equal to \( \sqrt{\pi}/2 \). \( f(r_c) \) is the force strength between particles at distance \( r_c \), the cutoff distance. \( g(r_c) \) is the radial pair distribution function at cutoff distance.

Our results for the elastic constants (in the solid range) are gathered in Fig. 3.11, where we have plotted the elastic constants \( C_{11}, C_{12} \) and \( C_{44} \) as functions of the temperature, as well as their fluctuation contributions. Whereas the constants themselves increase when the temperature is lowered, the opposite trend is observed for their fluctuation contributions. We can see that all the fluctuation terms go to 0 when the temperature decreases to 0. This result is obtained by making a linear extrapolation of our results at low temperatures. When \( T = 0 \), according to the formulas given in the previous section, the Born terms of the elastic constants are equal to the value of the elastic constants. The results from the linear extrapolation for the Born terms at low temperatures and the direct calculation at \( T = 0 \) are almost the same. Our numerical results are also displayed in Table 3.3.2. We have compared our results with some previous (few) calculations, for example in [1]. The agreement is excellent.

The shear modulus \( G \) can be computed by \( G = \mu - P = C_{44} - P \). From Fig. 3.12, we can see that the shear modulus of the system decreases linearly when temperature increases from 0.0 to 0.83, then it suddenly becomes zero at \( T = 0.84 \), which corresponds
Fig. 3.11 The elastic constants and the fluctuation terms of the elastic constants for LJ model system as a function of the temperature (in the solid region). The upper group of three (curves with hollow shapes) represent the total elastic constants, in decreasing order $C_{11}$, $C_{44}$ and $C_{12}$. The lower group of three (curves with solid shapes) represent the fluctuation parts, in decreasing order $C_{11f}$, $C_{12f}$ and $C_{44f}$. The pressure is $P = 1$. 

Elastic constants for LJ model system

$N = 1372, P \sim 1.0, T = 0.01-0.83$
Table 3.3.2 Elastic constants and the relevant fluctuation terms of elastic constants at
different temperatures for LJ system (LJ3d) in solid state (fcc crystal). We recall that
\( P = 1 \) throughout, and the melting temperature is \( T_m = 0.84 \).

<table>
<thead>
<tr>
<th>T</th>
<th>( C11_f )</th>
<th>( C12_f )</th>
<th>( C44_f )</th>
<th>( C11 )</th>
<th>( C12 )</th>
<th>( C44 )</th>
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<tbody>
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<td>-0.02</td>
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<td>11.74</td>
<td>18.22</td>
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</table>
Fig. 3.12 The shear modulus $G$ as a function of the temperature for LJ model system. The pressure is $P = 1$. Also plotted is the second “shear” modulus $G'$ defined by $G' = (C_{11} - C_{12})/2 - P$. 
Fig. 3.13 The bulk modulus $K$ as a function of the temperature for LJ model system. The pressure is $P = 1$.

to the phase transition temperature. This result is consistent with the behaviour of the specific volume. In an isotropic systems, a second formula for $\mu$ is $\mu' = (C_{11} - C_{12})/2$, giving a second formula for $G$, i.e. $G' = \mu' - P = (C_{11} - C_{12})/2 - P$. We show that above the phase transition temperature, for the LJ liquid, $G$ and $G'$ are both equal to zero. While below this temperature, the curves are quite distinct, with $G > 2G'$. This means that the LJ fcc crystal is quite anisotropic. The bulk modulus is calculated by $K = (C_{11} + 2C_{12} + P)/3$. We have plotted in Fig. 3.13 its evolution with the temperature, both at the solid and the liquid states. We can see that the bulk modulus for the system decreases almost linearly when the temperature increases. But the slope are quite different in each state, being strong in the solid, and quite weak in the liquid. At $T = 0$, we have $K \approx 80$, whereas at $T = 0.8$ (solid), $K \approx 20$. In the liquid side, $T > 0.84$, $K$ does not exceed 10. At the transition, $K$ is not well defined, so its numerical value in the plot may be accidental. Our numerical results are shown in Table 3.3.3. To summarize, we characterized the elastic properties of the LJ model
Table 3.3.3 Shear modulus $G$, bulk modulus $K$ and specific volume $v$ for LJ3d model system, as a function of the temperature. The “second” shear modulus, $G'$, is defined by $G' = (C_{11} - C_{12})/2 - P$. Results for $T = 0$ are extrapolations of $T > 0$ data. The pressure is $P = 1$ throughout, and the melting temperature is $T_m = 0.84$.

<table>
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<tr>
<th>$T$</th>
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<th>$G'$</th>
<th>$K$</th>
<th>$v$</th>
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<td>-0.15</td>
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<td>1.47</td>
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</table>
system using the stress fluctuation formalism. These results will serve as references when we tackle a more complex crystal, the AB13 superlattice.

### 3.3.3 AB13 model system

The superlattice formed by a binary mixture of hard spheres with a diameter ratio of $\sigma_B/\sigma_A = 0.58$, with the composition of one A-particle for 13 B-particles is a quite complex crystal [10]. The A particles form a simple cubic lattice. Inside each cube there is a centered icosahedra of 13 B particles. Because of a rotational freedom, each unit cell contains 8 cubes, that is 112 particles. This structure was found by experiments, theory and simulations of colloids, at high packing fraction. We use here this crystal as an example of complex crystals. In Figs. 3.14 and 3.15, the superlattice structure at $T = 0$ is shown.

![AB13 model system](image)

Fig. 3.14 Basic units of the primitive cell of the AB13 superlattice.

#### Simulation details

Again, in our molecular dynamics simulations, we use NPT and NVT ensembles. The potential between a pair of particles i and j, of species $\alpha$ and $\beta$, is

$$u_{\alpha\beta}(r_{i\alpha,j\beta}) = \begin{cases} 
4\epsilon_{\alpha\beta} \left[ \left( \frac{\sigma_{\alpha\beta}}{r_{i\alpha,j\beta}} \right)^{12} - \left( \frac{\sigma_{\alpha\beta}}{r_{i\alpha,j\beta}} \right)^{6} \right] & 0 < r_{i\alpha,j\beta} < r_{0\alpha\beta} \\
0, & r_{i\alpha,j\beta} > r_{0\alpha\beta}
\end{cases} \quad (3.13)$$

where $r_{0\alpha\beta} = 2^{1/6}\sigma_{\alpha\beta}$ is the position of the minimum of the LJ pair potential. Thus the pair potential is purely repulsive. Both the potential and its first derivative are zero.
at the cutoff. Therefore no impulsive correction is needed here. We present the pair potential in Fig. 3.16. The units correspond to the large spheres. We suppose the same mass for both species $m_A = m_B = 1$, $\epsilon_{AA} = \epsilon_{BB} = \epsilon_{AB} = 1$. $\sigma_{AA} = 1$, $\sigma_{BB} = 0.58$ and $\sigma_{AB} = (\sigma_{AA} + \sigma_{BB})/2 = 0.79$. The particle numbers are $N_A = 64$, $N_B = 13N_A = 832$, so the total number is $N = 896$. This potential is qualitatively close to hard sphere potentials. But it is continuous, allowing normal MD simulations with the LAMMPS code. The system is dense, under a pressure of $P = 114$ in order to stabilize the crystal phase.

We follow the same procedure as the case of LJ system. Here the starting temperature is $T = 1$ (solid). It is cooled down to $T = 0$, and heated up to $T = 3.5$. Between 0 and 3.5, several temperatures have been examined (see Table 3.3.4). The pressure was always fixed at the same value. So the heating and cooling are done using the NPT ensemble (rate $10^{-5}$). The equilibration is done using NPT as well. The statistics are gathered using the NVT ensemble.

### 3.3.4 Numerical results

We present the results in the same order as the LJ system.

First, we look at the specific volume. From the graph Fig. 3.17, we can see that there is a jump point along the curve $v(T)$, which implies the phase transition from crystal to liquid for AB13 model system. The melting temperature is about $T_m = 2.1$. 

---

Fig. 3.15 The primitive cell of the AB13 superlattice. (a) Front view. (b) Side view.
Fig. 3.16 Potentials for AB13 model system. Species 1 stands for A particles, 2 for B particles. Distances are measured by units of $\sigma_A$, potentials by $\epsilon_{AA}$.

Table 3.3.4 AB13 system. Temperature points studied by cooling or heating a reference equilibrium system at $T = 1.0$ (crystal). The pressure is $P = 114$ throughout.

<table>
<thead>
<tr>
<th>Process</th>
<th>Temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cooling</td>
<td>1.0  0.8  0.6  0.4  0.2  0.1  0.05</td>
</tr>
<tr>
<td>Heating</td>
<td>1.0  1.2  1.4  1.6  1.8  2.0  2.02</td>
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<tr>
<td></td>
<td>2.02  2.04  2.06  2.08  2.1  2.2  2.3</td>
</tr>
<tr>
<td></td>
<td>2.3  2.4  2.5  3.0  3.5</td>
</tr>
</tbody>
</table>
Fig. 3.17 Specific volume \((v = V/N)\) as a function of temperature for AB13 system. The pressure is fixed at \(P = 114\). All units refer to the large spheres.
Next, we calculate the RDF for a few temperatures. Fig. 3.18 shows the RDFs at very low temperature, $T = 0.1$. We see that the curves of different pairs have pronounced peaks, and they subsist at long distances. This means that the system is well structured (low temperature crystal).

In graph Fig. 3.19, the radial distribution functions for two higher temperatures are displayed. When the temperature is 2.0, the number of peaks reduces and the peaks become wider and lower than the $T = 0.1$ case, which reflects some disorder in the system. But still, the RDFs are crystal-like. So we are close to the melting. When the temperature is 2.1, the radial distribution functions of A particle and B particle become about 1 at long range, which is the feature of a liquid. As a result, the system undergoes the melting phase transition at this temperature. This is consistent with the specific volume result.

Now we characterize the elastic constants for the solid state. As before, we have three independent constants, the system being a cubic superlattice. From the graph Fig. 3.20, we can see that all the elastic constants for AB13 model decrease linearly when the temperature increases. Their values are much higher than the LJ system, because of the very high pressure here. On the other hand, we have, as previously, $C_{12}$
Fig. 3.19 Radial distribution functions of AB13 system at two different temperatures (2.0, 2.1).

close to $C_{44}$, and $C_{11}$ about two to three times larger.

Now, it is interesting to examine the fluctuation terms. From the graph Fig. 3.21, we see that these fluctuation terms almost increase linearly when the temperature increases. Remarkably, their values don’t go to zero when the temperature reaches zero. This is at odds with the situation of the fcc crystal. For $C_{11}$ and $C_{44}$, these fluctuations are about 50, whereas for $C_{12}$ it is actually negative! This shows the importance of taking the fluctuations terms into account, when computing the elastic constants, even at $T = 0$. In Table 3.3.5, detailed results for the AB13 solid are gathered.

The shear modulus is of course examined. We recall that $G = C_{44} - P$, and the alternative shear modulus is $G' = (C_{11} - C_{12})/2 - P$. From Fig. 3.22, we see that the shear modulus for AB13 system decreases when temperature increases from zero to 2.0. Then it becomes zero when temperature is above 2.1, which means that the melting transition happens at this temperature. This is again consistent with the specific volume behaviour. If we compare $G$ and $G'$, we see that they are very close, with $G$ slightly larger. This shows that the AB13 crystal is much more isotropic than
Fig. 3.20 Elastic constants vs. temperature for AB13 crystal.

Fig. 3.21 Elastic constants fluctuation terms vs. temperature for AB13 crystal.
Table 3.3.5 Elastic constants and fluctuation terms of elastic constants at different temperatures for AB13 model system in solid state (AB13 superlattice). The pressure is $P = 114$. The melting temperature is $T_m = 2.1$.

<table>
<thead>
<tr>
<th>T</th>
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<th>$C12_f$</th>
<th>$C44_f$</th>
<th>$C11$</th>
<th>$C12$</th>
<th>$C44$</th>
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</table>
Shear modulus as a function of the temperature for AB13 system, in both liquid and solid states. The pressure is fixed at $P = 114$. $G$ is the real modulus, $G'$ comes from an alternative definition $G' = (C_{11} - C_{12})/2 - P$.

The last elastic quantity shown is the bulk modulus $K$. From the graph Fig. 3.23, we can see that the bulk modulus for the AB13 system decreases when the temperature increases. There exists a singularity when temperature is about 2.1 (transition temperature). Again, the slope in the solid is much larger than the one in the liquid. At $T = 0$, this modulus is very important, $K \approx 1000$, about ten times the pressure. Near the transition, it is about 400, roughly four times the pressure. We also notice very little variation of $K$ just below and just above $T_m$. This should be related to the more isotropic feature of the solid (than the fcc crystal). Numerical results for these quantities are gathered in Table 3.3.6.

To summarize, our comparison of simple and complex crystals shows a difference in behaviour of the fluctuation terms at low temperatures. We also detected a more isotropic character of the AB13 superlattice. To our knowledge, the present results on
Table 3.3.6 Shear modulus $G$, bulk modulus $K$, system volume $V$ and box length $L$ for AB13 model system for $0 \leq T \leq 3.5$. We recall that $N = 896$, $P = 114$, $T_m = 2.1$, and $G' = (C_{11} - C_{12})/2 - P$. Results at $T = 0$ are extrapolations of the data for $T > 0$.

<table>
<thead>
<tr>
<th>$T$</th>
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<th>$G'$</th>
<th>$K$</th>
<th>$V$</th>
<th>$L$</th>
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Fig. 3.23 Bulk modulus vs. temperature for AB13 model

the elastic constants of AB13 crystal are the first ones. They will be completed with time-dependent properties displayed in the next chapter.
Chapter 4

Time-dependent properties

4.1 Introduction

In this chapter we investigate time-dependent properties. Some of them are numerical issues, such as the sampling time dependence of the physical quantities we compute, especially the stress fluctuations. Others are the intrinsic dynamical properties of the systems investigated, for example the shear-stress autocorrelation function $C(t)$, the shear-stress mean square displacement $h(t)$, the shear-stress relaxation modulus $G(t)$, the relationship between them, and their behaviour with the thermodynamic conditions. Another interesting property is the dynamic moduli, which can be obtained by sine and cosine transforms of $G(t)$. These are important mechanical properties of materials that are accessible to the experiments. We shall show that they can also be computed quite accurately and efficiently by our equilibrium MD simulations. Most of the theoretical formalism is based on ref.[16–18, 40]. Our investigations are focused on four systems: monodisperse two-dimensional Lennard-Jones (LJ2d) system, KA2d system, monodisperse three-dimensional Lennard-Jones (LJ3d) system and the AB13 system.

4.2 Sampling-time dependence of the stress fluctuations

In the stress fluctuation formalism, the static shear modulus $G$ is calculated according to

$$G = \mu_A - \mu_F$$

(4.1)
with \( \mu_A \) being the affine modulus, \( \mu_F \) the fluctuation of the stress. We have, by definition

\[
\mu_F = \beta V \langle (\hat{\tau} - \langle \hat{\tau} \rangle)^2 \rangle = \beta V \langle \hat{\tau}^2 \rangle - \beta V \langle \hat{\tau} \rangle^2 = \tilde{\mu}_F - \mu_*
\]

(4.2)

where \( \tilde{\mu}_F = \beta V \langle \hat{\tau}^2 \rangle \), and \( \mu_* = \beta V \langle \hat{\tau} \rangle^2 = \beta V \tau^2 \). Now we show that \( \mu_F \) depends on the sampling time used. In order to explain this and simplify the notations, we define a physical observable \( a \), with instantaneous values \( \hat{a} \). If \( a \) is measured over a time window \( \Delta t \), meaning \( n \) records at interval \( \delta t \), then we can write the time-average of \( a \) as

\[
\bar{a} = \frac{1}{n} \sum_{i=1}^{n} \hat{a}_i
\]

(4.3)

where \( \hat{a}_i = \hat{a}(t_i) \) with \( t_i = (i - 1)\delta t \) for \( i \) ranging from 1 to \( n \). If we have \( m \) trajectories of length \( n \), then the expectation value of \( a \) is the ensemble average given by

\[
E(a) = \langle \bar{a} \rangle = \frac{1}{m} \sum_{j=1}^{m} \bar{\hat{a}}_j
\]

(4.4)

where \( \bar{\hat{a}}_j \) denotes the time-average resulting from the number-\( j \) trajectory. For many observables, the ensemble and the time averages commute. They are called “simple averages”. Examples are the pressure \( P \), the average shear stress \( \tau \), the affine shear constant \( \mu_A \), and the mean value of \( \hat{\tau}^2 \) denoted by \( \tilde{\mu}_F \). On the other hand, the fluctuations of the observables do depend on \( \Delta t \). They are thus not simple averages. To show this, we imagine a long simulation run, consisting on \( N \) times \( \delta t \). This trajectory can be divided into \( m \) sub-trajectories each of length \( \Delta t \). The expectation value for the fluctuations of \( a \) is

\[
Fl(a) = \langle \bar{a}^2 - \langle a \rangle^2 \rangle = \frac{1}{m} \sum_{j=1}^{m} (\bar{\hat{a}}_j^2 - \bar{\hat{a}}_j^2).
\]

(4.5)

Obviously, \( Fl(a) \) will depend on \( \Delta t \) (because the second term in eq. (4.5) depends on \( \Delta t \)). It is actually a monotonically increasing function of \( \Delta t \), starting from zero when \( \Delta t = 0 \). In practice, when we compute \( \mu_F \), we have \( \hat{a} = \sqrt{\beta V} \hat{\tau} \). Thus

\[
\mu_F(\Delta t) = \beta V \langle \hat{\tau}^2 - \langle \hat{\tau} \rangle^2 \rangle
\]

(4.6)

is an increasing function of \( \Delta t \) (see Fig. 4.1).
The stress fluctuations $\mu_F(\Delta t)$ as a function of the sampling time $\Delta t$ \textit{(LJ2d system)}.

Its statistical value can be estimated when $\mu_F(\Delta t)$ approaches its plateau-value asymptotically. The real relaxation time of the system can be defined by $t_*$ such that the plateau-value as close as 1%. Fig. 4.2 shows our results for LJ2d and KA2d systems. We can observe that the time to reach the plateau depends on the system, and on the thermodynamic state. For the LJ2d system, $t_*$ is quite short (about 100). For the KA2d system, a binary mixture, it is certainly longer (more than 500 for the temperatures investigated).

Our results for 3d systems are gathered in Figs. (4.3-4.4). In these systems, $t_*$ appears to be quite short, either for the crystal or the liquids states. For AB13, it is of the order 10, for LJ3d system, about 100. Here the mixture reaches the plateau faster than the one-component system. This may be explained by the important size-disparity between the spheres, so that the small one have more freedom to move and help relax the system faster.

From eq. (4.1) and eq. (4.6), we can write, for the shear modulus

$$G(\Delta t) = \mu_A - \mu_F(\Delta t) = \mu_{A,ex} - \mu_{F,ex}(\Delta t),$$

as a function of the sampling time $\Delta t$. Fig. 4.5 shows this behaviour. It is seen that the shear modulus decays monotonously as a function of the sampling time for all
Fig. 4.2 The stress fluctuations $\mu_F(\Delta t)$ as a function of the sampling time $\Delta t$ (LJ2d and KA2d systems).

Fig. 4.3 The stress fluctuations $\mu_F(\Delta t)$ as a function of the sampling time $\Delta t$ (LJ3d system).

cases considered. Ultimately, however, all data sets level off as emphasized by the dashed horizontal lines. As may be seen from Fig. 4.5, much larger sampling times are needed for reaching the plateau of the shear modulus for the KA2d model, especially
around $T_g$, than for the LJ2d model. One thus has to check whether sufficiently large trajectories have been used to determine reliable long-time estimates. Please note that this has apriori nothing to do with an insufficient equilibration or ageing of the systems, but stems from the fact that the stress-fluctuations simply need time to explore the available phase space. The elastic moduli $G(T)$ and $K(T)$ presented previously thus stem from reasonable plateau values as one expects.

4.3 Shear-stress autocorrelation function

We study the shear-stress auto-correlation function (SACF) of the shear stress, defined by, as in[16],

$$C(t) = \beta V \langle \delta \dot{\tau}(t) \delta \dot{\tau}(0) \rangle.$$  \hspace{1cm} (4.8)

This function is important. It allows a better theoretical understanding of the dynamics in liquids[81], it leads to measurable quantities, such as the viscosity[14, 30], and to the shear modulus [40].

Two comments on $C(t)$ can be made. The first is the way we compute this cor-
Fig. 4.5 Sampling-time-dependence of the shear modulus $G_F(\Delta t) = \mu_{A,ex} - \bar{\mu}_{F,ex}(\Delta t)$ as a function of the sampling time $\Delta t$ for LJ2d and KA2d models and different temperatures as indicated. In all cases we reach a horizontal plateau $G(T)$ for sufficiently large $\Delta t$ as indicated by the dashed horizontal lines. (Here $\bar{\mu}_{F,ex}(\Delta t)$ is defined by eq. (4.6), using only the excess shear stress, and $\mu_{A,ex} = \mu_A - \rho k_B T$).

relation function. It is important to remind that the time-translational invariance is always assumed. This means that we have

$$C(t) = \beta V \left\langle \delta \hat{\tau}(t + t_0) \delta \hat{\tau}(t_0) \right\rangle,$$

where the horizontal bar denotes the gliding average over the time-origins $t_0$ taken within the trajectory. The second comment remarks that $C(t)$ is a correlation of the stress fluctuations. Thus it depends on the sampling time $\Delta t$. We can designate it as $C(t, \Delta t)$. What we expect is that if $\Delta t < t_*$ ($t_*$ is a typical relaxation time), then $C(t)$ is not correctly evaluated. In practice, we divide a long trajectory into $m$ sub-trajectories of $\Delta t$, ($j = 1, \ldots, m$). Thus the function $C(t)$ obtained from the $jth$ sub-trajectory is

$$C_j(t) = \beta V \bar{\delta \hat{\tau}(t + t_0) \delta \hat{\tau}(t_0)}_j = \beta V \bar{\hat{\tau}(t + t_0) \hat{\tau}(t_0)}_j - \mu_{*j}$$

where $\mu_* = \beta V \bar{\hat{\tau}}^2$ is obviously $\Delta t$-dependent, since this time-average is taken within
the time-interval $\Delta t$. The final result $C(t)$ is the ensemble average of $C_j(t)$:

$$
C(t) = \frac{1}{m} \sum_{j=1}^{m} C_j(t) = \frac{1}{m} \sum_{j=1}^{m} \left( \tilde{C}_j(t) - \mu_{*j} \right)
$$

(4.11)

where $\tilde{C}(t) = \beta V \bar{\tau}(t + t_0) \bar{\tau}(t_0)$ is the unshifted shear-stress SACF. At given $\Delta t$, we can commute the time and ensemble averages for $\tilde{C}(t)$, it is thus a simple average.

**Details for $\tilde{C}(t)$.** We denote $f(t) = \sqrt{\beta V} \bar{\tau}(t)$, the correlation time $t_k = k\delta t$, the correlation function $F(k)$. The time-average in $j$-th sub-trajectory is

$$
F(k)_j = \frac{\sum_i f(t_{i,j})f(t_{i+k,j}) + \ldots + f(t_{n-k,j})f(t_{n,j})}{(n - k)} \quad (0 \leq k \leq n - 1)
$$

(4.12)

and the final correlation function is, by ensemble average,

$$
F(k) = \frac{1}{m} \sum_j F(k)_j = \frac{1}{m} \sum_j \sum_i f(t_{i,j})f(t_{i+k,j})/[(n - k) * m],
$$

(4.13)

as the sums commute. $\tilde{C}$ is indeed a simple average. On the other hand, its precision can depend on $\Delta t$. To show this, we suppose that we start with $\Delta t_0$, corresponding to $n_0$ data points, coming from one of the $m_0$ sub-trajectories of a long run of $N$ records. Thus, we have

$$
F_0(k) = \frac{\sum_j \sum_i f(t_{i,j})f(t_{i+k,j})}{(n_0 - k) * m_0}
$$

(4.14)

If we set now $m = m_0 * n_s$ segments, each having $n = n_0 * n_s$ data points, then the new estimate of $F(k)$ is

$$
F_s(k) = \frac{\sum_j \sum_i f(t_{i,j})f(t_{i+k,j})}{(n - k) * m} = \frac{\sum_j \sum_i f(t_{i,j})f(t_{i+k,j})}{[(n_0 - k)/n_s) * m_0]}
$$

(4.15)

Obviously, except for $k = 0$, we have $F_s(k)$ different of $F_0(k)$. But the difference can be numerically neglectable, if we ensure that for each $k$, there are at least $10^2$ data for the statistics, between the time and ensemble averages.

On the other hand, $\mu_{*}$ is not a simple average. We have, for a given $\Delta t$,

$$
\mu_{*} = \frac{\sum_j \mu_{*j}}{m} = \sum_j \left( \sum_i f(t_{i,j})/n \right)^2 /m
$$

(4.16)

showing that the sum over $i$ ($n$ terms) and the one over $j$ ($m$ terms) are not com-

mutable. Furthermore, $\mu_{*}$ depends strongly on $\Delta t$, being a maximum for $\Delta t = 1.\delta t$, and decreasing to its plateau value for $n >> 1$. 84
From the above discussion, it is clear that the SACF, for a finite $\Delta t$, should be noted $C(t, \Delta t)$. However, we can drop the second argument if $\Delta t$ is large enough ($\Delta t \gg t_*)$, after the function has converged.

Fig. 4.6 - 4.9 show results on this topic. We see that indeed, they all depend on $\Delta t$. But the convergence can be reached after a reasonable sampling time, of the order $\Delta t \approx 100$, for the state points explored. Another observation is that clearly, the crystals (LJ2d, LJ3d and AB13) show strong non monotonic SACF at short times, whereas in the liquid, the function is a monotonic decreasing function of time. In the low temperature glass (KA2d), the function is also monotonic, although less smooth as in the liquid. Another general feature is that the function goes to zero at long times, for all systems and all states. This means that it is impossible to predict the shear modulus of a solid from this limit, as some authors believe [28].

![Fig. 4.6 The shear-stress autocorrelation function (SACF) of the LJ2d system as a function of the time, for several sampling times $\Delta t$. The left panel corresponds to a low temperature crystal, $T = 0.2$, the right one to a liquid, $T = 0.8$.](image)

The sequence $\mu_F(\Delta t)$ can be generated by $C(t)$, as shown in [16]. We have (for more details, see the next section),

$$\frac{\mu_F(\Delta t)}{\mu_F} = 1 - \frac{2}{\Delta t^2} \int_0^{\Delta t} dt (\Delta t - t) C(t) \equiv 1 - y(\Delta t)$$  \hspace{1cm} (4.17)

We can thus define a time $\theta$ such that $\mu_F(\theta)$ is very close to $\mu_F$, for example at 95%, or $y(\theta) = 0.05$. 

85
Fig. 4.7 The SACF of the KA2d system as a function of the time, for several sampling times $\Delta t$. The left panel corresponds to a low temperature glass, $T = 0.1$, the right one to a liquid, $T = 0.8$.

Fig. 4.8 The SACF of the LJ3d system as a function of the time, for several sampling times $\Delta t$. The left panel corresponds to a low temperature crystal, $T = 0.2$, the right one to a liquid, $T = 0.8$.

In Fig. 4.10, the SACF $C(t)$ and the function $y(\Delta t)$ of the monodisperse LJ2d system are presented, at the liquid and solid states, respectively. We see that for both states, the $C(t)$ curves decay to 0 quite quickly, in about one time unit. The “real” relaxation time of the SACF, as suggested in ref[40], could be defined by the time for which $y(\Delta t)$ reaches 0.05 (the blue line in the inset), corresponding here to about 5 time units for the states explored. The functions, however, are much longer lasting in the glass (the KA2d model), near $T_g$, as shown in Fig. 4.11. A good estimate of
Fig. 4.9 The SACF of the AB13 system as a function of the time, for several sampling times $\Delta t$. The left panel corresponds to the AB13 superlattice, $T = 2.0$, the right one to a liquid, $T = 2.2$.

The SACF relaxation time can again be obtained by the method suggested in ref[40]. Using this definition, we obtained results displayed in Table 4.1, where we see indeed that the relaxation in the glass (near $T_g$) is about 500, compared to (about) 5 in the crystal.

Table 4.1 Various relaxation times associated with the stress auto-correlation function $C(t)$. $t_C$ corresponds to $C(t_C)/C(0) = e^{-1}$, $t_{C1}$ to $C(t_C)/C(0) = 0.05$, and $\theta$ to $y(\theta) = 0.05$.

<table>
<thead>
<tr>
<th>System</th>
<th>LJ2d (liquid)</th>
<th>LJ2d (crystal)</th>
<th>KA2d (liquid)</th>
<th>KA2d (glass)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T$</td>
<td>0.7</td>
<td>0.6</td>
<td>0.4</td>
<td>0.3</td>
</tr>
<tr>
<td>$t_C$</td>
<td>0.1</td>
<td>0.1</td>
<td>0.2</td>
<td>0.3</td>
</tr>
<tr>
<td>$t_{C1}$</td>
<td>0.6</td>
<td>0.5</td>
<td>5.4</td>
<td>99</td>
</tr>
<tr>
<td>$\theta$</td>
<td>6.4</td>
<td>5.4</td>
<td>32.5</td>
<td>500</td>
</tr>
</tbody>
</table>
Fig. 4.10 The SACF \( C(t) \) and the function \( y(\Delta t) \) (inset) for the monodisperse LJ2d system. The red (dashed) curve corresponds to a crystal \( (T = 0.6) \), while the black (continuous) curve represents a liquid state \( (T = 0.7) \). The blue dash-dotted line in the inset indicates the level \( y = 0.05 \).

4.4 The shear-stress mean-square displacement

In [17], we introduced the "mean square displacement" (MSD) of the shear-stress, defined by

\[
h(t) = \beta V \left\langle \left( \tilde{\tau}(t) - \tilde{\tau}(0) \right)^2 \right\rangle / 2.
\]  

(4.18)

It is straightforward to show that

\[
h(t) = \tilde{C}(0) - \tilde{C}(t) = C(0) - C(t)
\]  

(4.19)

As \( \tilde{C}(0) = \tilde{\mu}_F \) and \( C(0) = \mu_F \), we can invert eq. (4.19) to write

\[
\tilde{C}(t) = \tilde{\mu}_F - h(t); \quad C(t) = \mu_F - h(t).
\]  

(4.20)

Now we show that \( h(t) \) is a simple average. We have, for a given \( \Delta t \),

\[
h(t, \Delta t) = \beta V \left\langle \left( \tilde{\tau}(t + t_0) - \tilde{\tau}(t_0) \right)^2 \right\rangle / 2, \quad \text{for} \quad t \leq \Delta t
\]  

(4.21)
Fig. 4.11 The SACF $C(t)$ and the function $y(\Delta t)$ (inset) for the glass-forming KA2d system. The red (dashed) curve corresponds to a glass ($T = 0.3$), near $T_g = 0.32$, while the black (continuous) curve represents a liquid state ($T = 0.4$). For the liquid, $C(0) \approx 34$ (the upper black dot), while for the glass, $C(0) \approx 33$ (the lower black dot). The blue dash-dotted line in the inset indicates the level $y = 0.05$.

where the large bar denotes the gliding average over $t_0$ within the a $\Delta t$ package. In detail, for the $j$-th sub-trajectory, we have

$$h(k)_j = \left[ ((f(t_{1+k,j}) - f(t_{1,j}))^2 + ... + (f(t_{n,j}) - f(t_{n-k,j}))^2) / (n - k) \right] (0 \leq k \leq n - 1),$$

thus

$$h(k) = \frac{\sum_{j=1}^{m} h(k)_j}{m},$$

showing that the time-average and the ensemble average commute. On the other hand, similarly to the $\tilde{C}$ function, the numerical precision may depend on $\Delta t$ chosen. But this can be controlled ensuring enough statistics for each $k$ value, as mentioned previously.

Figs. 4.12 - 4.15 show examples of $h(t)$ function. We see that the function starts at 0, and saturates at long times to its plateau value. For all curves but the third one, the relaxation time $t_A$ is about 1. The evolution of $h(t)$ is much slower for the KA2d liquid and glass close to the glass transition temperature, and is oscillatory for the low temperature AB13 crystal. The first observation can be understood by longer stress
Fig. 4.12 The shear stress mean square displacement as a function of the time for the LJ2d model systems. The first time point is $t_1 = 0.005$.

Fig. 4.13 The shear stress mean square displacement as a function of the time for the KA2d model systems. Here, the first time point is $t_1 = 0.05$ (except for $T = 0.28$, where $t_1 = 0.1$).

relaxation times in the system close to the glass transition. The second case, quite intriguing, may be caused once again by the size disparity between the two species,
causing faster time scales for the small spheres.

While $h(t)$ reaches the asymptote rapidly at $t \approx t_A$, the sampling-time dependent stress fluctuation $\mu_F(\Delta t)$ reaches its plateau value $\mu_F$ much more slowly, at a time...
scale $\Delta t \approx t_\ast$. Remarkably, we can understand it by the relation between these two time-dependent properties. Starting from $\mu_F$, we have

$$
\mu_F(\Delta t) = \left\langle \frac{1}{n} \sum_i (a_i - \bar{a})^2 \right\rangle
= \left\langle \bar{a}^2 - \bar{a}^2 \right\rangle
= (1/2n^2) \left\langle \sum_i \sum_l (a_i - a_l)^2 \right\rangle
$$

(4.24)

where $\langle ... \rangle$ stands for the ensemble-average over the trajectories (of length $n$). The last line can be written as

$$
\mu_F(\Delta t) = (1/n^2) \left\langle \sum_{i=l+1}^{n} \sum_{l=1}^{n-l} (a_i - a_l)^2 \right\rangle
= (1/n^2) \left\langle \sum_{l=1}^{n-1} \sum_{k=1}^{n-l} (a_{l+k} - a_l)^2 \right\rangle
= (1/n^2) \sum_{k=1}^{n-1} \sum_{l=1}^{n-l-k} \langle (a_{l+k} - a_l)^2 \rangle
= (1/n^2) \sum_{k=0}^{n-1} (n-k) g(k)
$$

(4.25)

where “mean-square advance” $g(k) = \langle (a_{l+k} - a_l)^2 \rangle$ is independent of $l$ (time translational invariance). Noting $g(k) = 2h(k)$ (the MSD of the shear stress) and supposing $\delta t << 1$, we can write eq. (4.25) into an integral:

$$
\mu_F(\Delta t) = (2/\Delta t^2) \int_0^{\Delta t} dt (\Delta t - t) h(t)
$$

(4.26)

eq (4.27)

eq. (4.26) can give us a fair idea of $t_\ast$, knowing $t_A$ of the function $h(t)$. For this, we define $y(\Delta t) = 1 - \mu_F(\Delta t)/\mu_F$, where $\mu_F$ is the plateau value of $\mu_F(\Delta t)$. According to eq. (4.26), we have

$$
y(\Delta t) = 1 - (2/\Delta t^2) \int_0^{\Delta t} dt (\Delta t - t) h(t)/\mu_F.
$$

(4.27)

Assuming $t_\ast$ is such that $y(t_\ast) = 1\%$, and $t_\ast >> t_A$, from eq. (4.27), we obtain,

$$
0.01 \approx t_A/t_\ast
$$

(4.28)
i.e. \( t_\ast \approx 100t_A \). So we have now an order of magnitude of \( t_\ast \). As \( C(t) = C(0) - h(t) \).

For a given \( \Delta t \) this implies \( C(t, \Delta t) = \mu_F(\Delta t) - h(t) \), i.e.,

\[
C(t, \Delta t) = (2/\Delta t^2) \int_0^{\Delta t} dt (\Delta t - t)h(t) - h(t).
\]

(4.29)

Thus we have a converged \( C(t) \) only if \( \Delta t >> t_\ast \). This fact is important, especially when we want to compute the shear stress relaxation modulus \( G(t) \), as we shall see in the next section.

### 4.5 The shear relaxation modulus

The shear relaxation modulus, \( G(t) \), is a central rheological property of materials. It is defined by the linear response macroscopic relation

\[
G(t) = \frac{\partial \tau(t; \gamma)}{\partial \gamma}
\]

(4.30)

where \( \gamma \) is a small step strain imposed at \( t = 0 \), and \( \tau(t) \) is the averaged stress increment of the system. \( G(t) \) gives the measure of the viscoelasticity of a system. It is important that we can compute this function by using statistical physics tools. Previously, we showed that the static shear modulus \( G \) can be computed using the stress fluctuations formalism, by Monte Carlo (MC) or molecular dynamics (MD) simulations. This quantity is denoted by \( G_{eq} \) here, to distinguish it from \( G(t) \). Obviously, \( \lim_{t \to \infty} G(t) = G_{eq} \). For an elastic body, \( G_{eq} > 0 \), and for a fluid, \( G_{eq} = 0 \). Given that \( \tau(t) = \langle \hat{\tau}(t) \rangle \), what is the relation between \( G(t) \) and the SACF \( C(t) \), or \( \tilde{C}(t) \). Many authors assumed \( G(t) = \tilde{C}(t) = C(t) \). We shall demonstrate that this is only true for the fluids, where we have \( \tau = 0 \) (at equilibrium), and \( G_{eq} = 0 \). In more general case, we start from

\[
\tau(t) = \int_{-\infty}^{t} ds G(t - s) d\gamma(s)/ds
\]

\[
= G(t - s)\gamma(s)|_{-\infty}^{t} - \int_{-\infty}^{t} ds dG(t - s)/ds \gamma(s)
\]

(4.31)
where we have used integration by parts. Here we suppose $\gamma(t)$ a step function ($\gamma(t) = \gamma$ for $t \geq 0$ and $\gamma(t) = 0$ for $t < 0$),

$$\tau(t) = G(0)\gamma - \int_0^t \chi(s)ds \cdot \gamma$$

$$= \left[ G_{eq} + \int_t^\infty \chi(s)ds \right] \cdot \gamma$$

(4.32)

where, in the first line, we define $\chi(t) \equiv -G'(t) = G'(-t)$ is the response function (or the "after-effect function", and in the second line, $G_{eq}$ appears naturally as an integration constant. As the response function is related to $C(t)$ by $\chi(t) = -C'(t)$ (see [14]), we obtain

$$G(t) = C(t) + G_{eq}$$

(4.33)

or

$$G(t) = \tilde{C}(t) + \mu_A - \tilde{\mu}_F.$$  

(4.34)

Furthermore, from eq. (4.33), we see that $G(t)$ can be related to the stress MSD in a very simple way:

$$G(t) = \mu_A - h(t).$$

(4.35)

Eq. (4.35) shows that $G(t)$ can be evaluated from a simple averages $\mu_A$ and $h(t)$. As $h(t)$ relaxes much faster than $\mu_F(\Delta t)$, eq. (4.35), $G(t)$ relaxes actually much faster than $C(t)$ itself, reaching $G_{eq}$ faster than $G_F(\Delta t) = \mu_A - \mu_F(\Delta t)$. It also shows clearly that $G(0) = \mu_A$, the affine shear elasticity, which is always positive, either the system is solid, or even liquid.

The Figs. 4.16 - 4.19 show the function $G(t)$ as obtained from eq. (4.35) for our model systems. The common feature is that all the curves shown (except one) reach their limit values quite rapidly, at a time scale of 1. The exception concerns the low temperature AB13 crystal, which oscillates for some time before being stabilized. This is of course for the same reason as its $h(t)$ curve.

To summarize, eq. (4.35) not only gives the function $G(t)$ in a quite accurate way (simple averages), it also provides the equilibrium shear modulus without the necessity of long sampling times $\Delta t$. So it is an advantageous way of studying the elasticity and the viscoelasticity by simulations. However, some tests have to be done using eq. (4.34) with converged $C(t)$. The resulting $G(t)$ seems quite comparable.
Fig. 4.16 The shear-stress relaxation modulus $G(t)$ of the LJ2d system for several temperatures.

Fig. 4.17 The shear-stress relaxation modulus $G(t)$ of the KA2d system for several temperatures.
Fig. 4.18 The shear-stress relaxation modulus $G(t)$ of the LJ3d system for several temperatures.

Fig. 4.19 The shear-stress relaxation modulus $G(t)$ of the AB13 system for several temperatures.
4.6 Dynamic moduli

4.6.1 Definitions and simple models

In a simulation, the function $G(t)$ can be collected from the dynamics of the system. This quantity can also be measured in rheological experiments. However, more frequently, people can use oscillatory stresses to measure the dynamic modulus $G^*$ (see refs. [12, 13]). This quantity is frequency dependent $G^* = G^*(\omega)$ with $\omega$ the angular frequency. More precisely, if we impose an oscillatory shear strain $(\gamma_0 \sin(\omega t))$, given, in complex representation, by

$$\gamma(t) = \gamma_0 \exp(i\omega t), \quad (4.36)$$

in the linear response regime the average shear stress will be

$$\tau(t) = G^*(\omega)\gamma(t) \equiv (G''(\omega) + iG''(\omega))\gamma_0 \exp(i\omega t), \quad (4.37)$$

The real part of $G^*$, $G'(\omega)$, is the storage modulus, corresponding to the part of the response of the system in phase with $\gamma(t)$, i.e. the elastic response; the imaginary part $G^*$, $G''(\omega)$, is the loss modulus, corresponding to the part that is dephased $\pi/2$ w.r.t. $\gamma(t)$, i.e. the viscous response.

The physical meaning of $G''$ can be shown by computing the average power that the stresses should provide to maintain the periodic oscillation of the system. We write, per unit volume, this average as

$$\text{Power}_T = \frac{1}{T} \int_0^T \tau(t) * \dot{\gamma}(t) dt, \quad (4.38)$$

where $T = 1/2\pi \omega$ is the period of the solicitation $\gamma(t)$. Using $\gamma(t) = \gamma_0 \sin(\omega t)$ and $\tau(t) = Im(G^* \gamma_0 \exp(i\omega t)) = G' \sin(\omega t) + G'' \cos(\omega t)$, we obtain $\text{Power}_T = G''.\gamma_0.\omega/2 \propto G''$. As this is the power lost by the system, $G''$ is indeed the “loss modulus”.

In order to relate $G^*$ with $G(t)$, let’s consider first a viscoelastic liquid, with $G_{eq} = 0$, and compute $G^*_{l}$ ($l$ for “liquid”). We start from the Boltzmann superposition principle [13], which gives

$$\tau(t) = \int_{-\infty}^t G_l(t - t') \dot{\gamma}(t') dt'. \quad (4.39)$$
From eq. (4.36), we obtain $\dot{\gamma}(t) = i\omega_0 \gamma(t)$. After a change of the variable in eq. (4.39), by posing $s = t - t'$, we obtain

$$
\tau(t) = i\omega_0 \gamma(t) \int_0^\infty G_l(s) \exp(-i\omega s) ds = \gamma(t) \omega \int_0^\infty G_l(s) (\sin(\omega s) + i \cos(\omega s)) ds \tag{4.40}
$$

By identification, we obtain

$$
G_l'(\omega) = \omega \int_0^\infty G_l(t) \sin(\omega t) dt \\
G_l''(\omega) = \omega \int_0^\infty G_l(t) \cos(\omega t) dt. \tag{4.41}
$$

In general situations where $G_{eq} \neq 0$, eq. (4.41) is generalized to

$$
G'(\omega) = G_{eq} + \omega \int_0^\infty (G(t) - G_{eq}) \sin(\omega t) dt \\
G''(\omega) = \omega \int_0^\infty (G(t) - G_{eq}) \cos(\omega t) dt. \tag{4.42}
$$

This shows that $G'$ corresponds to the Fourier-sine transform of $G(t)$, and $G''$ to its Fourier-cosine transform. Their low frequency limits are well known:

$$
\lim_{\omega \to 0} G'(\omega) = G_{eq} \\
\lim_{\omega \to 0} G''(\omega)/\omega = \int_0^\infty (G(t) - G_{eq}) dt = \int_0^\infty C(t) dt = \eta \tag{4.43}
$$

where we used eq. (4.33), and the Green-Kubo relation to obtain viscosity coefficient $\eta$ (see [14]) . For a liquid, $\eta$ is the usual viscosity. For a solid, $\eta$ represents the slope of $G''$ for $\omega \to 0$.

We can also draw some conclusions on the high frequency limit. Supposing, for the stress ACF $C(t)$, $\lim_{t \to \infty} C(t) = 0$, and using $C(0) = \mu_F$, from eq. (4.42) we can deduce (by integration by parts),

$$
\lim_{\omega \to \infty} G'(\omega) = \mu_A = G(0) \\
\lim_{\omega \to \infty} G''(\omega) = 0. \tag{4.44}
$$

This tells us that the system, either in liquid or solid states, possesses a high frequency non-zero elasticity, given by the affine modulus $\mu_A$. On the other hand, $G''$ vanishes at high frequencies.
Now we can compare these general trends with the ones obtained from two simple classical rheological models, the Maxwell model and the Kelvin-Voigt (K-V) model [13]. They consist of the association of two elementary rheological elements, the elastic element, a spring of constant $G$, and a viscous element of viscosity $\eta$. The Maxwell model is a serial association and the K-V model a parallel association (see Fig. 4.20).

\[ \sigma = G\gamma \quad \text{(the index E stands for elastic),} \]

\[ \sigma = \eta \dot{\gamma} \quad \text{(index V for viscous).} \]

For the Maxwell model, we have

\[ \sigma = \sigma_E = \sigma_V, \gamma = \gamma_E = \gamma_V, \text{ thus the rheological equation is:} \]

\[ \frac{\dot{\sigma}}{G} + \frac{\sigma}{\eta} = \dot{\gamma}. \quad (4.45) \]

To determine $G(t)$, we can make a "relaxation" experiment: we set

\[ \gamma(t) = \begin{cases} 0 & \text{if } t < 0, \\ \gamma_0 & \text{if } t \geq 0, \end{cases} \quad (4.46) \]

and calculate $\sigma(t)$ for $t > 0$ according to eq. (4.45), given that at $t = 0^+$, it is the elastic element which can react instantly: $\sigma(0^+) = G\gamma_E(0^+) = G\gamma(0^+) = G\gamma_0$. The solution is

\[ G(t) = Ge^{-t/\tau_M} \quad (t \geq 0) \]

with $\tau_M = \eta/G$ the Maxwell relaxation time. This result corresponds to a viscoelastic fluid: $G_{eq} = \lim_{t \to \infty} G = 0$, and $G(0) = G$. 

Fig. 4.20 Rheological models. $\sigma$ is the shear stress, $\gamma$ is the shear strain.
The dynamic moduli can be obtained in two ways: a) by solving directly eq. (4.45), using \( \gamma(t) = \gamma_0 e^{i\omega t} \) and \( \sigma(t) = G^* \gamma(t) \); b) by eq. (4.42). Both give

\[
G'(\omega) = G \frac{(\omega \tau_M)^2}{1 + (\omega \tau_M)^2}; \quad G''(\omega) = G \frac{\omega \tau_M}{1 + (\omega \tau_M)^2}.
\] (4.48)

We see that \( G'(0) = G''(0) = 0 \), \( \lim_{\omega \to 0} G''(\omega) = G' \tau_M = \eta \), \( \lim_{\omega \to \infty} G'(\omega) = G \), \( \lim_{\omega \to \infty} G''(\omega) = 0 \). For \( \omega < \tau_M^{-1} \), \( G' > G'' \) (the system is more viscous at low frequencies), for \( \omega > \tau_M^{-1} \), \( G' < G'' \) (the system is more elastic at high frequencies), at \( \omega = \tau_M^{-1} \), there is a crossover, \( G' = G'' \), thus \( \tau_M^{-1} \) is the characteristic frequency of the system.

Fig. 4.21 shows \( G(t) \) and \( G^* \) of the Maxwell model.

![Figure 4.21 Rheological moduli of the Maxwell model, \( \tau_M = \eta/G \).](image)

Fig. 4.21 Rheological moduli of the Maxwell model, \( \tau_M = \eta/G \).

For the K-V model, we have \( \gamma = \gamma_E = \gamma_V \), \( \sigma = \sigma_E + \sigma_V \), thus the rheological equation is:

\[
G \gamma + \eta \dot{\gamma} = \sigma \] (4.49)

To determine \( G(t) \), we can again make the “relaxation” experiment, by solving eq. (4.49) for \( t > 0 \), where \( \dot{\gamma} = 0 \). The solution is

\[
G(t) = G; \quad (t > 0)
\] (4.50)

This result corresponds to a solid: \( G_{eq} = \lim_{t \to \infty} G \). For this model, \( G(0) \) is unphysical, because it is singular (looking at eq. (4.49) ).

The dynamic moduli can be obtained by solving directly eq. (4.49), using \( \gamma(t) = \gamma_0 e^{i\omega t} \) and \( \sigma(t) = G^* \gamma(t) \). This gives

\[
G'(\omega) = G; \quad G''(\omega) = G \omega \tau_M \] (4.51)
Obviously, $G''$ is unphysical at high frequencies, because it tends to infinity. By using eq. (4.42), we can complete the singular part of $G(t)$, to obtain

$$G(t) = G\tau_M \delta(t) + G = \eta \delta(t) + G. \quad (4.52)$$

From eq. (4.52), we have the stress ACF, $C(t=0) \to \infty$, in contradiction to $C(0) = \mu_F$ (finite) for any real system.

Fig. 4.22 shows $G(t)$ and $G^*$ of the K-V model.

Another measurable mechanical quantity is the compliance $J^*$ (see for example ref. [13]), defined by $\gamma(t) = J^*(\omega)\sigma(t)$, where $\sigma(t) = \sigma_0 e^{i\omega t}$ is imposed. Clearly $J^* = 1/G^*$. It is customary to write $J = J' - iJ''$. For the Maxwell model, we have

$$J'(\omega) = \frac{1}{G}; \quad J''(\omega) = \frac{1}{G\omega\tau_M}. \quad (4.53)$$

Again, we see that for $\omega < \tau_M^{-1}$, $J' < J''$ (the viscosity dominates at low frequencies), for $\omega > \tau_M^{-1}$, $J' > J''$ (the elasticity dominates at high frequencies).

Fig. 4.23a shows $J^*$ of the Maxwell model.

As for the compliance, we obtain, for the K-V model,

$$J'(\omega) = \frac{1}{G(1 + (\omega\tau_M)^2)}; \quad J''(\omega) = \frac{\omega\tau_M}{G(1 + (\omega\tau_M)^2)}. \quad (4.54)$$

Fig. 4.23b shows $J^*$ of the K-V model. One can see that the $J^*$ moduli display better (than $G^*$) the viscoelastic character of the solid represented by this model.
Fig. 4.23 The compliance of the Maxwell and Kelvin-Voigt models, $\tau_M = \eta/G$.

From the simulation data $G'(\omega)$ and $G''(\omega)$, the expressions of $J'$ and $J''$ are

$$J'(\omega) = \frac{G'(\omega)}{G'(\omega)^2 + G''(\omega)^2}, \quad J''(\omega) = \frac{G''(\omega)}{G'(\omega)^2 + G''(\omega)^2}. \quad (4.55)$$

These relations will be useful for us to convert our $G^*$ moduli to $J^*$ ones. Their limits are obvious: $J'(0) = 1/G_{eq}$, $J'(\infty) = 1/\mu_A$, and $J''(0) = J''(\infty) = 0$. But $J''$ is not zero for all $\omega$ (see the next sub-section). As $J'(0)$ must be finite, we shall only discuss this quantity when our system is in solid state.

**4.6.2 Simulation results**

Turning to our systems, it is an easy task to calculate $G^*$ once $G(t)$ is known. It would be interesting to investigate how our $G^*$ compare with those predicted by the simple rheological models. For the somewhat delocalization high frequency limit, we use Filon’s method [30] to perform the sine and cosine transformations. Our results are gathered in Figs. 4.24 - 4.31. The general features for the liquids seem to follow qualitatively the Maxwell model, if we identify $G$ in the Maxwell model with $\mu_A$, the high-frequency limit of our storage modulus $G'$. A “Maxwell time” ($\tau_M$) can be estimated from the maximum of $G''(\omega)$ in our liquids. Once we have $G$ and $\tau_M$, theoretical fit can be done using eq. (4.48). The results are shown in Figs. 4.25, 4.27, 4.29 and 4.31. The overall agreement is remarkable. However, two differences can be noted. The first is a peak of $G'$ at some intermediate frequency (resonance), absent in the Maxwell model. The second difference is that for $\omega > 1/\tau_M$, our $G''$ seem to
decay faster than the Maxwell model, for all four liquid systems. These differences show the limitation of such a simple rheological model, even for the liquid state. When our systems are solids, the Maxwell model naturally fails, because the model gives $\lim_{\omega \to 0} G'(\omega) = 0$ always, instead of $G_{eq} > 0$. And the K-V model does not apply to our solids either, because, clearly we have two limit values for the storage modulus ($G_{eq}$ and $\mu_A$) instead of only one in the K-V model (see Fig. 4.22). Furthermore, our $G''$ are very different from $G''$ by the K-V model, which is unphysical for high frequencies. Nevertheless, we will (later) compare the compliance of our solids with the K-V model, to see whether there is some agreement for certain frequency zones. Another point about the dynamic moduli of our solids is that the loss modulus $G''$ is not at all negligible for some frequencies. Its maximum can even be as important as $G_{eq}$. This shows the potential importance of this quantity (representing the dissipation), even for a solid. It highlights certainly the interest of our method, allowing accurate computation of this quantity. Furthermore, we can see that $G''(\omega)$ displays more complex features for KA2d and AB13 solids than the LJ2d and LJ3d solids. This can certainly be traced back to the former systems being binary mixtures, rather than monodisperse systems, as the LJ2d and LJ3d systems.
Fig. 4.24 Dynamic moduli of LJ2d model in the triangular crystal structure $T = 0.6$. Two recording time intervals for the stress tensor are used to generate $G(t)$, i.e. $5 \times 10^{-3}$ (circles) and $2 \times 10^{-4}$ (squares). This allows to cover both the low frequency and the high frequency zones of the dynamic moduli.

Fig. 4.25 Dynamic moduli of LJ2d model system in liquid state $T = 0.8$. The fit uses $\tau_M = 0.10, G = \mu_A$ and eq. (4.48).
Fig. 4.26 Dynamic moduli of KA2d model system as a low temperature glass $T = 0.1$.

Fig. 4.27 Dynamic moduli of KA2d model system in liquid state $T = 0.8$. The fit uses $\tau_M = 0.091$, $G = \mu_A$ and eq. (4.48).
Fig. 4.28 Dynamic moduli of LJ3d model system in crystal state $T = 0.8$.

Fig. 4.29 Dynamic moduli of LJ3d model system in liquid state $T = 0.9$. The fit uses $\tau_M = 0.088$, $G = \mu_A$ and eq. (4.48).
The fit uses $\tau_M = 0.031$, $G = \mu_A$ and eq. (4.48).

Fig. 4.30 Dynamic moduli of AB13 model system in crystal state $T = 2.0$.

Fig. 4.31 Dynamic moduli of AB13 model system in liquid state $T = 2.2$. The fit uses $\tau_M = 0.031$, $G = \mu_A$ and eq. (4.48).
Now it’s interesting to plot the compliance of our systems in the solid state, in order to compare them with the predictions of the K-V model. They are displayed in Figs. 4.32 - 4.35. Again, the time parameter $\tau_M$ can be extracted from the simulation $J''$ curve, at its maximum, i.e. $\tau_M = 1/\omega_{max}$, where $\omega_{max}$ is the frequency for the maximum of $J''$. This give $\tau_M = 0.22, 0.33, 0.14$ and $0.063$ for LJ2d, KA2d, LJ3d and AB13 systems respectively. As for $G$ to be used in eq. (4.54), we have the choice of $G_{eq}$, or $\mu_A$, with $G_{eq}$ be the natural choice. The fit using $G_{eq}$ is satisfactory for the mixtures (KA2d and AB13), for $J''$ overall and $J'$ in the range $\omega < \omega_{max}$. Whereas for the one-component systems, $J''$ is too large. If we use $\mu_A$ then $J''$ is well fitted, but $J'$ must be shifted by $1/\mu_A$. These general trends could depend on temperature. Further investigations are needed.
Fig. 4.33 Compliance moduli of the KA2d model system, at $T = 0.1$ (glass). The fit corresponds to $\tau_M = 0.33$ and $G = G_{eq}$ in eq. (4.54).

Fig. 4.34 Compliance moduli of the LJ3d model system, at $T = 0.8$ (crystal). Two sets of fit are performed with the K-V model. Both using the time $\tau_M = 0.14$. For $J'_1$ and $J''_1$, $G = G_{eq}$ is used. For $J'_2$ and $J''_2$, $G = \mu_A$ is used, and $J'_2$ is shifted by $1/\mu_A$. 
Fig. 4.35 Compliance moduli of the AB13 model system, at $T = 2.0$ (AB13 superlatice). The fit (with the K-V model) corresponds to $\tau_M = 0.063$ and $G = G_{eq}$.

4.7 Topical conclusions

To summarize, we presented in this chapter general considerations on time dependent properties. We first examined the sampling-time dependence of the stress fluctuations. Then we studied the shear stress autocorrelation function $C(t)$, the shear stress mean square displacement $h(t)$, the shear stress relaxation modulus $G(t)$, and the relations between them. The key relation $G(t) = \mu_A - h(t)$ is used to calculate the shear stress relaxation modulus for our model systems at liquid and solid states. The resulting curves show satisfactory accuracy, and allow for the computation of the dynamic moduli. This quantity, of high experimental relevance, is shown to be now available by equilibrium MD simulations.
Chapter 5

Conclusions

5.1 Summary

In this thesis, we have studied elasticity, viscoelasticity, and glass transition of model systems by computer simulations, using equilibrium stress and volume fluctuations. The model systems considered are as follow: monodisperse hard sphere fcc crystal, polydisperse hard sphere fcc crystal, a glass-former consisting of the two dimensional Kob-Andersen model (KA2d system), its companion system, a monodisperse LJ2d system. Then the monodisperse LJ3d system and a binary repulsive mixture AB13 system, forming respectively simple fcc and complex superlattice crystals, are studied. From these systems we obtained some new and interesting results.

First, as a simple test of the relation linking the bulk modulus and the volume fluctuations under constant pressure, we computed the bulk modulus of (slightly) polydisperse hard sphere crystals at high pressure by Monte Carlo NPT simulations. We showed that the equation of state proposed by Bartlett is not valid for high pressures, whereas our results give the effects of the polydispersity on the average density and the bulk modulus. Furthermore, the volume distribution displays, as expected, a Gaussian histogram, indicating good statistics of our simulations. This approach is a relatively simple way to calculate the bulk modulus of general systems. It is thus potentially interesting for various purposes (testing liquid theories, studying systems near a phase transition, or under high pressure, etc).

Second, we investigated the two dimensional 80-20 Kob-Andersen model under constant moderate pressure with focus on the elastic moduli calculation. Our results
show that the KA2d model is a reasonably good glass-former. We examined three ways of determining the glass transition temperature $T_g$, and found that they give consistent results. The shear modulus as a function of the temperature $G(T)$ has been characterized. The result has been found to depend strongly on sampling time, with very slow convergence near the glass transition. We observed a continuous cusp-type variation of $G(T)$ across the glass transition. This finding is in qualitative agreement with recent numerical work using similar glass-forming colloidal systems in two and three dimensions [70], but in contradiction with predictions from the mode-coupling theory.

Along with the KA2d model, we also investigated a simple companion model provided by monodisperse 2dLJ beads, which crystallizes at low temperature. In contrast with the KA2d model, we observed a discontinuous jump of $G(T)$ at the freezing transition.

Third, we investigated the shear stress fluctuations in simple and complex crystals including Lennard-Jones (LJ3d) model systems and a binary AB13 model systems. For each system, the pressure is fixed. The evolution of the systems with temperature is examined. We were able to locate the melting transition temperature by the discontinuous jump of the elastic constants. This jump is consistent with the jump in specific volume at the transition. In the low temperature region, the fluctuation terms of elastic constants for these two model systems are scrutinized. While for the simple fcc crystal, these terms go to zero for $T = 0$, they are notably non-zero for the more complex AB13 crystal, contributing to about 10% for the shear modulus. This is less than in a glass, but still significant. The degree of anisotropy of the two crystals can be examined by looking at the two ways of computing the shear modulus, using $G = C_{66} - P$, or $G_2 = (C_{11} - C_{12})/2 - P$, by recalling that they are equal for an isotropic system. We observe that the fcc crystal is quite anisotropic, with $G \approx 3G_2$.

On the other hand, the AB13 superlattice seems to be much more isotropic, since we have in this case $G \approx G_2$, with $G$ only slightly above $G_2$ at very low temperatures.

Fourth, we examined the time dependent properties of our four model systems such as the sampling time dependence of the stress fluctuations, the shear-stress autocorrelation function $C(t)$, the shear stress mean square displacement $h(t)$, the shear-stress relaxation modulus $G(t)$, the relationship between these functions, and their behaviour with the thermodynamic conditions. The dynamic moduli, $G'(\omega)$ and $G''(\omega)$, and the
compliance moduli $J'(\omega)$ and $J''(\omega)$, are also computed. From the sampling time $\Delta t$ dependence of the stress fluctuations, we can see that the sampling time more than 100 in reduced units is sufficient for the LJ2d system, the AB13 system and the LJ3d system, while it is more than 1000 for KA2d system at least, when the temperature is close to $T_g$. Clearly, this glassy system (near $T_g$) evolves much more slowly than crystal systems. As for the shear stress autocorrelation function $C(t; \Delta t)$, we have confirmed, as expected, the sampling time dependence of this function. We note that in most cases, $\Delta t \approx 100$ gives convergent $C(t)$. For the shear stress mean square displacement $h(t)$, the AB13 system show oscillation at low temperature during the relaxation time. The function is seen to be non-monotonic in the crystal, at low temperatures. This feature has not been observed in the permanent elastic bodies [17]. The long-time plateau value of $h(t)$ increases with the increment of temperature at solid state while the trend is the opposite at liquid state. This is not surprising since it corresponds to $\mu_F$. We note that $h(t)$ is smoother than $C(t)$, being a simple average. In general, it indeed is a better function for the calculation of $G(t)$ via $G(t) = \mu_A - h(t)$, as pointed out in [17]. Here, for some state points, we have computed shear relaxation modulus $G(t)$. The function displays a rapid decrease from $G(0) = \mu_A$ (the affine elasticity), and it reaches its equilibrium modulus $G_{eq} = \mu_A - \mu_F$ at long times. From $G(t)$, we have computed the dynamic moduli, and compared our results with the Maxwell and the Kelvin-Voigt rheological models. The four systems studied have qualitatively the same behaviour as the Maxwell rheological model when they are liquid (with some small differences discussed in Chapter 4). At solid state, the AB13 system and the KA2d system show more complicated features than the other two systems. We speculate that the potentials for the AB13 system and the KA2d system are more complex and they have two kinds of particles so that they have different response to the external fields. Here, for the storage modulus $G'$, clearly the Maxwell model does not apply since we have two distinct limit $G'$ values, i.e. $G'_0 = G_{eq}$ for $\omega = 0$ and $G'_\infty = \mu_A$ for $\omega \rightarrow \infty$, with $G'_\infty > G'_0$. Because of these two values, the Kelvin-Voigt model does not apply either. Interestingly, we have again a peak of $G'$, at some intermediate frequency, where $G'_{\text{max}} > \mu_A$ (resonance). As for the loss modulus $G''$, while it is zero for low and high frequencies, as expected, its value is by no means neglectable for some intermedi-
ate frequencies, with its maximum even comparable in magnitude to $G_{eq}$. This shows that even in these simple model systems, the energy dissipation in the solid state can be important for some frequencies. The chapter ends with a brief discussion on the compliance moduli $J'$ and $J''$ of our models in solid state.

5.2 Outlook

Many future extensions of our work are possible:

Concerning the elasticity, we can use the elastic constants to probe solid-solid phase transitions under high pressure, or with temperature. More general potentials, such as three-body, or EAM potentials should be examined. The viscoelastic functions, $G(t)$, $G'$ and $G''$ can be calculated for many soft matter systems and, by comparison with experimental results, allow a better modelling of these systems. For the glass transition, our study of the KA2d system must be carried further. For example, the trajectory analysis can be implemented. More independent configurations should be used, in order to reduce statistical errors in our moduli, especially near the glass transition. In parallel, it would be interesting to obtain similar expressions as $G(t) = \mu_A - h(t)$ for hard sphere systems, since these systems are objects of many theoretical and experimental investigations (hard sphere colloids).
Bibliography


[56] P. Cicuta, E. J. Stancik, and G. G. Fuller. Shearing or compressing a soft glass
in 2d: time-concentration superposition. Physical review letters, 90(23):236101,
2003.

in binary two-dimensional colloidal suspensions. Physical review letters, 97(7):
078301, 2006.

[58] F. Ebert, P. Dillmann, G. Maret, and P. Keim. The experimental realization of a
two-dimensional colloidal model system. Review of Scientific Instruments, 80(8):
083902, 2009.

[59] C. L. Klix, G. Maret, and P. Keim. Discontinuous shear modulus determines the

ments of growing amorphous order and non-monotonic dynamic correlations in a

[61] L. Berthier and G. Tarjus. The role of attractive forces in viscous liquids. The

glass-forming liquids in two, three, and four dimensions. Physical review letters,

length scales in supercooled liquids on approaching the glass transition. The Journal

[64] G. M. Hocky and D. R. Reichman. A small subset of normal modes mimics the
properties of dynamical heterogeneity in a model supercooled liquid. The Journal


Résumé de la thèse en français

0.1 Cadre général

L’élasticité et la viscoélasticité sont des propriétés essentielles de beaucoup de matériaux, en état solide ou fluides complexes. Il est important de les connaître pour de nombreuses applications pratiques. Ces grandeurs sont sensibles aux conditions thermodynamiques, notamment la température et la pression. En outre, leur brusque changement est souvent la signature d’une transition de phase. Ces aspects montrent qu’il est souhaitable de pouvoir étudier ces propriétés par simulation sur ordinateur, afin de modéliser les matériaux, et de prédire leur comportement dans des conditions potentiellement difficiles d’accès par expérience. Dans cette thèse, l’élasticité, viscoélasticité et la transition vitreuse de liquides vitrifiants et de solides modèles sont étudiés par des simulations moléculaires à l’équilibre en utilisant le formalisme de fluctuation de contraintes. Ce formalisme permet des études de l’élasticité et de la viscoélasticité sans déformer le système, mais en utilisant les fluctuations des contraintes à l’équilibre. Bien que proposé à la fin des années 1960 [1], il n’a réellement attiré l’attention des chercheurs du domaine qu’à la fin des années 1980 [2,4]. Et certaines avancées théoriques sont survenues assez récemment (dans les années 2000)[5,8,18]. Notre objectif est d’appliquer ce formalisme à des systèmes modèles représentatifs de systèmes colloïdaux ou atomiques pour répondre à des questions d’ordre fondamental, surtout liées à des transitions de phase. Les systèmes modèles étudiés sont composés de systèmes de Lennard-Jones à deux et trois dimensions, un mélange binaire de Lennard-Jones (LJ) des particules constituant la version 2d du modèle bien connu de Kob-Anderson (modèle KA2d), un mélange binaire AB13 de particules répulsives et un mélange ternaire de sphères dures. Nous étudions principalement l’effet de la température $T$ sur les propriétés mécaniques
Les systèmes, en particulier les transitions de phase telles que la cristallisation ou la transition vitreuse. Les résultats sont présentés en deux parties, à savoir les propriétés statiques et les propriétés dépendantes du temps. La plupart des résultats sont obtenus avec le code LAMMPS, de simulation en dynamique moléculaire à équilibre.

Les propriétés statiques que nous avons étudiées sont, pour les systèmes isotropes (verres et liquides), le module de cisaillement $G$ et le module de compression $K$, et pour les cristaux, les constantes élastiques. Ce sont des grandeurs sensibles à une transition de phase (solidification). Ainsi, nous examinons leur évolution en fonction de la température, et leur comportement à une transition de phase. Ainsi, nous avons montré une nette différence de comportement du module de cisaillement lors d’une transition vitreuse et lors d’une cristallisation. Alors que dans le 2ème cas, il y a une nette discontinuité de $G$, dans le 1er, l’évolution semble être continue. Un autre aspect intéressant est la contribution de la partie non-affine à l’élasticité, qui est calculable avec notre formalisme de fluctuation des contraintes (via le terme correspondant à ces fluctuations). Ce que nous voulons examiner est d’une part l’évolution de cette contribution en fonction de la température, et d’autre part son importance à température nulle. Nos résultats montrent que pour le cristal simple (du type cfc), ce terme est nul à $T = 0$, comme attendu, par contre, il est non nul pour un super-réseau du type AB13, contribuant jusqu’à 10% de la valeur du module élastique. Pour un verre, naturellement, ce terme est encore plus important (environ la moitié du module).

Dans la partie consacrée aux propriétés dépendantes du temps, nous avons étudié plusieurs aspects et fonctions. Tout d’abord, nous avons exploré la dépendance du temps d’échantillonnage de nos résultats statiques, et montré que cette influence est forte pour les fluctuations de contrainte. Deuxièmement, la dynamique d’équilibre est étudiée par le biais de l’autocorrélation des contraintes de cisaillement et déplacement quadratique moyen de la contrainte, ainsi que la relation entre ces deux fonctions temporelles. L’objet principal, cependant, est le module de relaxation de la contrainte de cisaillement $G(t)$ et les modules dynamique associée $G'(\omega)$ et $G''(\omega)$, avec $\omega$ la fréquence. Ces fonctions caractérisent la viscoélasticité de nos systèmes. En utilisant une formule proposée récemment (Wittmer et al, Phys. Rev. E 2016), nous avons pu déterminer ces fonctions avec précision et étudier leur comportement pour nos systèmes.
0.2 Thèmes étudiés et résultats

Tout d’abord, comme un simple test de la relation reliant le module de compression aux fluctuations de volume sous pression constante, nous avons calculé le module de compression des cristaux de sphères dures polydisperses ( légèrement) à haute pression par des simulations Monte Carlo dans l’ensemble \( NPT \) [82] (voir Figure 1). Nous avons montré que l’équation d’état proposée par Bartlett [49] n’est pas valable pour des pressions élevées. En plus, nos résultats donnent les effets de la polydispersité sur la densité moyenne et le module de compression. De plus, la distribution du volume affiche, comme prévu, un histogramme gaussien, indiquant de bonnes statistiques de nos simulations. Cette approche est un moyen relativement simple de calculer le module de compression des systèmes généraux. Il est donc potentiellement intéressant à diverses fins (test des théories des liquides, étude des systèmes près d’une transition de phase, ou sous haute pression, etc.).

Deuxièmement, nous avons étudié le modèle de Kob-Andersen à deux dimensions dans une composition 80-20 [83], sous une pression modérée constante en mettant
l’accent sur le calcul des modules élastiques. Nos résultats montrent que le modèle KA2d est un excellent système vitrifiant. Nous avons examiné trois façons de déterminer la température de transition vitreuse \( T_g \), et nous avons trouvé qu’ils donnent des résultats cohérents. Nous avons caractérisé \( G \) et \( K \) du modèle KA2d vitrifiant, et comparé les résultats avec un système de LJ2d monodisperse (voir Figure 2). Alors que ce dernier représente un saut significatif en \( G \) à la cristallisation, le système KA2d affiche un comportement de type continu à dérivée divergente à la transition vitreuse \( T_g \), suivant la loi \( G(T) \propto (1 - T/T_g)^a \) avec \( a = 0.6 \), confirmant qualitativement les résultats des simulations sur le système KA3d en 2013, mais en contradiction avec les prédictions basées sur la théorie de couplage de mode. Nous avons constaté que le résultat dépend fortement du temps d’échantillonnage, que la convergence est lente près de la transition vitreuse.

![Structure à basse température.](image1)

![Module de cisaillement vs \( T \).](image2)

Fig. 2 Structure et le module de cisaillement \( G \) pour le modèle KA2d.

Troisièmement, nous avons étudié les fluctuations de la contrainte de cisaillement dans les cristaux simples et complexes en examinant les systèmes modèles Lennard-Jones (LJ3d) et les systèmes binaires AB13. Pour chaque système, la pression est fixée. L’évolution des systèmes à température est examinée. Nous avons pu localiser la température de transition de fusion par le saut discontinu des constantes élastiques. Ce saut est cohérent avec le saut dans le volume spécifique à la transition. Dans la région basse température, les termes de fluctuation des constantes élastiques (élasticité non-affine) pour ces deux systèmes modèles sont examinés. Alors que pour le cristal cfc simple, ces termes vont à zéro pour \( T = 0 \), ils sont clairement non nuls pour le cristal plus complexe, à savoir AB13, contribuant à environ 10% pour le module de
cisaillement. Cette proportion est moins importante que dans un verre, mais encore significative (voir Figure 3).

(a) Constantes élastiques du solide AB13. (b) Termes de fluctuation du solide AB13.

Fig. 3 Constantes élastique et termes de fluctuation du solide AB13 (super-réseau) en fonction de la température.

Le degré d’anisotropie des deux cristaux peut être examiné en regardant deux manières de calculer le module de cisaillement, en utilisant \( G = C_{66} - P \), ou \( G' = (C_{11} - C_{12})/2 - P \), en rappelant qu’elles sont égales pour un système isotrope. Nous observons que le cristal cfc est tout à fait anisotrope, avec \( G \approx 3G' \). D’autre part, le super-réseau AB13 semble beaucoup plus isotrope, puisque nous avons dans ce cas \( G \approx G' \), avec \( G \) seulement légèrement au-dessus de \( G' \) à très basses températures (voir Figure 4).

Comme dernier thème, nous avons examiné les propriétés dépendantes du temps de nos systèmes modèles, telles que la dépendance au temps d’échantillonnage des fluctuations de contraintes, la fonction d’autocorrélation de la contrainte de cisaillement \( C(t) \), le déplacement carré moyen de la contrainte de cisaillement \( h(t) \), le module de relaxation de contrainte de cisaillement \( G(t) \), la relation entre ces fonctions et leur comportement avec les conditions thermodynamiques. Les modules dynamiques de conservation et de perte, ainsi que la complaisance sont également calculés. A partir de la dépendance des fluctuations de contraintes en fonction du temps d’échantillonnage \( \Delta t \), nous constatons que le temps d’échantillonnage supérieur à 100 (en unités réduites) est suffisant pour le système LJ2d, le système AB13 et le système LJ3d, alors qu’il
Fig. 4 Module de cisaillement $G$ en fonction de la température des solides LJ3d (cfc) et AB13 (super-réseau) respectivement. Deux modes de calcul sont représentés (voir texte).

Les modèles de calcul sont représentés (voir texte).

Le module de cisaillement pour le solide LJ3d est supérieur à 1000 pour KA2d, lorsque la température est proche de $T_g$. De toute évidence, ce système vitreux (près de $T_g$) évolue beaucoup plus lentement que les systèmes cristallins. Quant à la fonction d’autocorrélation de la contrainte de cisaillement $C(t, \Delta t)$, nous avons confirmé, comme prévu, la dépendance du temps d’échantillonnage de cette fonction. Nous notons que dans la plupart des cas, $\Delta t \approx 100$ est suffisant pour la convergence de $C(t)$. Pour le déplacement carré moyen $h(t)$, le système AB13 montre une oscillation à basse température pendant le temps de relaxation. La fonction se montre non monotone dans le cristal, à basses températures. Cette particularité n’a pas été observée dans les corps élastiques permanents [17]. La valeur du plateau à long terme de $h(t)$ augmente avec l’augmentation de la température à l’état solide tandis que la tendance est à l’opposé à l’état liquide. Cela n’est pas surprenant puisqu’il correspond à la fluctuation des contraintes de cisaillement $\mu_F$.

Ensuite, pour certains états thermodynamiques, nous avons calculé le module de relaxation de cisaillement $G(t)$. Comme indiqué dans [17], cette fonction peut se calculer via $G(t) = \mu_A - h(t)$, où $\mu_A$ est le module de l’élasticité affine. La fonction affiche une diminution rapide de $G(0) = \mu_A$, et il atteint son module d’équilibre $G_{eq} = \mu_A - \mu_F$ à au temps long. A partir de $G(t)$, nous avons calculé les modules dynamiques, et avons comparé nos résultats aux modèles rhéologiques simples de Maxwell et de Kelvin-
Voigt. Les quatre systèmes étudiés ont qualitativement le même comportement que le modèle de Maxwell quand ils sont liquides (avec quelques petites différences discutées au chapitre 4). À l’état solide, le système AB13 et le système KA2d présentent des caractéristiques plus compliquées que les deux autres systèmes, qui sont monodisperses. Nous supposons cela peut s’expliquer par le caractère binaire de KA2d et AB13. Pour nos solides, concernant le module de conservation $G'$, il est clair que le modèle Maxwell ne s’applique pas puisque nous avons deux valeurs limites de $G'$ non nulles (à basse et à haute fréquences). En raison de ces deux valeurs, le modèle de Kelvin-Voigt ne s’applique pas non plus. Fait intéressant, nous avons de nouveau un pic de $G'$, à une fréquence intermédiaire, où $G'_{\text{max}} > \mu_A$ (résonance). Quant au module de perte $G''$, alors qu’il est nul pour les fréquences basses et hautes, comme prévu, sa valeur n’est nullement négligeable pour certaines fréquences intermédiaires, son maximum est parfois même comparable en ordre de grandeur à $G_{\text{eq}}$. Cela montre que même dans ces systèmes de modèles simples, la dissipation d’énergie à l’état solide peut être importante pour certaines fréquences. Le chapitre se termine par une brève discussion sur les modules de complaisance de nos systèmes à l’état solide (voir Figure 5).

Fig. 5 Les modules dynamiques du modèle KA2d en état liquide et solide respectivement. $G'(\omega)$ est le module de conservation, $G''(\omega)$ est le module de perte.
0.3 Perspectives

De nombreuses extensions futures de notre travail sont possibles:

Concernant l’élasticité, nous pouvons utiliser nos méthodes de calcul des constantes élastiques pour étudier les transitions de phase solide-solide sous haute pression, ou en fonction de la température. Des potentiels plus généraux, tels que des potentiels à trois corps ou du type EAM devraient être examinée, afin d’étudier les propriétés élastiques de matériaux plus réalistes, surtout en fonction de la température et de la pression. Les fonctions viscoélastiques peuvent être calculées pour de nombreux systèmes de la matière molle, pour des comparaisons avec les résultats expérimentaux, permettent une meilleure modélisation de ces systèmes. Pour la transition vitreuse, notre étude du système KA2d doit être poursuivie. Par exemple, l’analyse des trajectoires peut être mise en œuvre. Plus de configurations indépendantes devraient être utilisées, afin de réduire les erreurs statistiques dans nos modules, en particulier près de la transition vitreuse. En parallèle, il serait intéressant d’obtenir des expressions similaires comme $G(t) = \mu_A - h(t)$ pour les systèmes de sphères dures, puisque ces systèmes sont des modèles théoriques et expérimentaux (colloïdes de sphères dures).