Shock waves in an augmented one-dimensional atom chain

G STOLTZ†‡
†CERMICS, Ecole Nationale des Ponts et Chaussées, 77455 Marne-la-Vallée, France
‡CEA/DAM Ile-de-France, BP 12, 91680 Bruyères-le-Châtel, France
E-mail: stoltz@cermics.enpc.fr

Abstract. We derive here a simplified discrete one-dimensional (1D) model describing some important features of shock waves. In order to avoid expensive multidimensional simulations, 1D models are commonly used, but the existing ones often exhibit some spurious physically irrelevant behavior. Here we build a 1D model with perturbations arising from mean higher-dimensional behavior. The coupling of the system with a deterministic heat bath in the Kac-Zwanzig fashion allows us to derive a generalized Langevin equation for the system, without a priori fixing the temperature in the shocked region. This deterministic problem with several degrees of freedom is then reduced to a simpler stochastic problem with memory. Some numerical results are provided, that illustrate and confirm the qualitative correctness of the model.

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1. Introduction

The aim of this study is to derive and assess the validity of a simplified microscopic model of shock waves that could help to calibrate parameters for macroscopic descriptions. Shock waves are intrinsically propagative phenomena. It is thus reasonable to describe them within a 1D macroscopic theory. In some cases depending on the geometry, this approximation has proven to be correct [3].

A 1D lattice seems an appropriate model that could, in addition, allow for some mathematical treatment and thus a better theoretical understanding of the phenomena and mechanisms at play. Indeed, many mathematical results are known about the behavior of waves in 1D lattices, concerning the existence of localized waves [10, 26], the form of those waves in the high-energy limit [8] or in the low-energy limit [9], or the behavior under shock [6]. There also exist extended results for a particular interaction between sites, the Toda potential [27]: the structure of a 1D shock is then precisely known, at least in some regime [23].

We begin in Section 2 with some introduction to 1D lattice motion, and briefly report on some theoretical results and numerical experiments on piston-impacted shocks. It is shown that, in the absence of a specific treatment, the shock profiles generated significantly differ from shock waves. Especially, their thicknesses grow linearly with time [17, 23], there is no usual equilibration downstream the shock front [4, 19, 23], and relaxation waves do not behave as expected. Indeed, one would expect the shock wave to be a self-similar jump separating two domains at local thermal equilibrium at different temperatures. The relaxation waves should then catch up the shock front and weaken the shock wave until it disappears. So, we have to introduce higher-dimensional effects, at least in an averaged way. This is performed in Section 3. The connection of the chain with a heat bath consisting of a large number of harmonic oscillators, seems to be a good remedy for spurious 1D effects. The shocks generated have constant thicknesses and relaxation waves appear to be properly modelled. We eventually present some simulation results in Section 4.

2. The pure 1D model

2.1. Description of the lattice model

Consider a one-dimensional chain of particles with nonlinear nearest-neighbor interactions, described by a potential $V$. Initially, the particles are at rest at positions $X_n(0) = nd$, which is an equilibrium state for the system. All the masses are set to 1. The normalized displacement of the $n$-th particle from its equilibrium position is $x_n(t) = \frac{1}{d}(X_n(t) - X_n(0))$. The following normalization conditions [17] for the interaction potential $V$ can be used:

$$V(0) = 0, \quad V'(0) = 0, \quad V''(0) = 1.$$  \hspace{1cm} (2.1)

The first condition is more a shift on the energy reference, the second one expresses the fact that $x = 0$ is the equilibrium position, and the last one amounts to a rescaling of time. The so-called ”reduced relative displacement” is defined as $\delta x_n(t) = x_{n+1}(t) - x_n(t)$. 
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The Hamiltonian of the system is:

$$H_S(\{q_n, p_n\}) = \sum_{n=-\infty}^{\infty} V(q_{n+1} - q_n) + \frac{1}{2}p_n^2,$$

(2.2)

where \((q_n, p_n) = (x_n, \dot{x}_n)\). The Newton equations of motion read:

$$\ddot{x}_n = V'(x_{n+1} - x_n) - V'(x_n - x_{n-1}).$$

(2.3)

The potential taken here can either have a physical origin, like the 1D Lennard-Jones potential:

$$V_{LJ}(x) = \frac{1}{8} \left( \frac{1}{(1+x)^4} - \frac{2}{(1+x)^2} \right),$$

(2.4)

or more mathematical motivations, like the one-parameter Toda potential [27]:

$$V_{Toda}^b(x) = \frac{1}{b^2} (e^{-bx} - 1 + bx).$$

(2.5)

Define \(b = -V''(0)\). The parameter \(b\) measures at the first order the anharmonicity of the system. For the Lennard-Jones potential \(b = 9\), and for the Toda potential, the parameter \(b\) introduced in the definition (2.5) is indeed equal to \(-\frac{d^3V^b}{dx^3}(0)\).

2.2. Shock waves in the 1D lattice

2.2.1. A brief review of the existing mathematical and numerical results

A shock can be generated using a ”piston”: the first particle is considered as being of infinite mass and constantly moving at velocity \(u_p\). We refer to [5] for a pioneering study of those shocks in 1D lattices, to [15, 17, 19] for careful numerical experiments and formal analysis, and to [23] for a rigorous mathematical study in the Toda case. All of these studies identify the parameter \(a = bu_p\) as critical. When \(a < 2\), the velocity of the downstream particles converge to the piston velocity, in analogy with the behavior of a harmonic lattice (see Figure 1). When \(a > 2\), the particles behind the shock experience an oscillatory motion (see Figure 2). This behavior is quite similar to what is happening in hard-rod fluids (see [19] for a more precise description of that phenomenon), and has to be linked to the exchange of momenta happening when two particles collide in a 1D setting. This was also noticed for other potentials such as the Lennard-Jones potential, and can be used to define specific 1D thermodynamical averages [4].

In the case of a strong shock \((a > 2)\) and in the Toda case, the displacement pattern is particularly well understood from a mathematical point of view [23]: the lattice can be decomposed in three regions. In the first one, for \(n > c_{\text{max}}t\), the particles have ”almost” not felt the shock yet, and their displacements are exponentially small. The second region, whose thickness grows linearly in time \((c_{\text{min}}t < n < c_{\text{max}}t)\), is composed of a train of solitons. Recall that solitons are particular solutions of the Toda lattice model, and correspond to localized waves [27]. In the third region \((n < c_{\text{min}}t)\), the lattice motion converges to an oscillatory pattern of period 2 (binary wave). The motion behind the

\[‡\] Note that we use \(b = 2a\) with the notation of [17].
Figure 1. Relative displacement (left) and velocity profiles (right) versus particle index for a weak shock at a representative time: number of particles $N_{\text{part}} = 500$, Toda parameter $b = 1$, piston velocity $u_p = 0.2$, so that $a = 0.2$. The particle are taken initially at rest at their equilibrium positions.

Figure 2. Relative displacement (left) and velocity profiles (right) versus particle index for a strong shock at time $T = 100$: $b = 10$, $u_p = 1$, so that $a = 10$. The particles are initially at rest.

shock is asymptotically described by the evolution of a single oscillator (see [4] for a precise description of this behavior). There is no local thermal equilibrium in the usual sense (i.e. the distribution of the velocities is not of Boltzmann form). This was already mentioned in [19].

2.2.2. Density plots. To get a better understanding of the shock patterns, it is convenient to represent the system in terms of local density. This local density can be obtained as a function of the local average of the interatomic distances, both in space and time. We restrict ourselves to a local average in space.

More precisely, the local averaged interatomic distance of the $n$-th length is denoted by $\bar{\delta x}_n$, and given by:

$$\bar{\delta x}_n = \sum_{i=-\infty}^{+\infty} \alpha_j \delta x_{n+j}.$$
The local density \( \rho_n \) is then defined as:

\[
\rho_n = (1 + \delta x_n)^{-1}.
\]

The weights \( \{\alpha_j\} \) are chosen in practice to be non negative and of sum equal to one. A convenient choice is for example:

\[
\alpha_j = C^{-1} \cos \left( \frac{j}{2M+1} \pi \right)
\]

for \(-M \leq j \leq M\), and \(\alpha_j = 0\) otherwise. The constant \(C\) is a normalization factor:

\[
C = \sum_{j=-M}^{M} \cos \left( \frac{j}{2M+1} \pi \right).
\]

The integer \(M\) is the local range of averaging.

Figure 3 gives the densities corresponding to the relative displacement patterns of Figures 1 and 2.

\[\text{Figure 3. Density patterns for the relative displacement pattern of the weak shock of Figure 1 (left) and the strong shock of Figure 2 (right). The local averaging range is } M = 50.\]

2.2.3. Simulation of piston compression

We first implement a preliminary thermalization. The particles are taken initially at rest at their equilibrium positions. We then generate displacements \(x_n\) and velocities \(\dot{x}_n\) from the probability density

\[
d\nu = \bigotimes_{n=-\infty}^{\infty} Z^{-1} e^{-\frac{1}{2} \beta_x (x_n^2 + \dot{x}_n^2)} \, dx_n \, d\dot{x}_n,
\]

with \(Z = 2\pi / \beta_x\). The initial displacements and velocities are then of order \(1 / \sqrt{\beta_x}\). Notice that we take small initial displacements, so we approximate the full potential \(V(x)\) by its harmonic part \(\frac{1}{2} x^2\). This approximation is of course justified only at the beginning of the simulation, when displacements are small enough. After this initial perturbation, we let the system free to evolve during a typical time \(T_{\text{init}} = 10\). The simulations were performed using a Velocity Verlet scheme, the time step being chosen to have a relative energy conservation \(\Delta E / E\) of about \(10^{-3}\).
At time $T_{\text{init}}$, the piston impact begins: the first particle is kept moving toward the right at constant velocity $u_p$.

Let us emphasize that the shock patterns are robust, in the sense that they remain essentially unchanged when initial thermal perturbations are supplied. This point was already noted in [19] where the authors gave numerical evidence of that fact. While rigorously proven only in the Toda lattice case for a lattice initially at rest at equilibrium, the above shock description seems then to remain qualitatively valid for a quite general class of potentials and with random initial conditions. A comparison of the different profiles is made in Figures 4 and 5. The profiles are indeed quite conserved, especially the density profiles.

**Figure 4.** Relative displacement profiles for a thermalized strong shock using a Toda potential with $b = 10$, and comparison with the reference profile corresponding to a lattice initially at rest. The piston speed is $u_p = 0.3$ (so that $a = 3$), $\sqrt{\beta}$ = 0.02.

**Figure 5.** Local density profiles corresponding to Figure 4 with $M = 50$. Dashed line: reference profile. Solid line: Thermalized profile. Notice that both patterns almost coincide.

For strong shocks ($a > 2$), the shock front thickens linearly with time as can be seen in Figure 6. This is in contradiction with what is observed in shock propagation experiments as well as in 3D numerical simulations. Moreover the velocity distribution behind the
shock front shows that the downstream particles experience a (quasi-)oscillatory motion in the range $[0, 2u_p]$. This is of course not the case for 3D simulations, where the particle velocities are much less correlated, and appears to be a pure 1D effect.

We emphasize once again that initial thermal perturbations are not sufficient to remedy these spurious 1D effects since the patterns obtained in Figures 4 and 5 are very similar. In the sequel we are going to build a 1D model that enables us to get rid of these undesired effects.

2.2.4. Simulation of relaxation waves In order to study the relaxation waves, the piston is removed after a compression time $t_0$, and the system evolves freely during time $t_1 - t_0$.

The results are once again not physically satisfactory. The soliton train of Figure 7, which was less visible in Figure 4, is not destroyed by the relaxation waves. It travels on and
widens since the solitons move away from each others (the distance between the fastest ones, that is, the more energetic ones, and the slowest ones, increases). We emphasize that the energy remains localized in those waves, so there is no damping of these solitons. Rarefaction is only observed in the region behind the soliton train.

On the other hand, in 3D simulations or in experiments, one observes a progressive damping of the whole compressive wave. This is a second spurious effect of the 1D model we would like to get rid of and that our model will able to deal with.

3. Introduction of mean higher-dimensional effects

The results of the previous Section indicate the need for a modeling of perturbations arising from the transverse degrees of freedom existing in higher dimensional simulations. Such perturbations will interfere with the shock front composed of a soliton train, and possibly damp this soliton train. Perturbations in the longitudinal direction, such as thermal initialization for the \( x_n \), cannot do this, as shown by Figures 4 and 5.

Actually, some facts are already known about the influence of 3D effects on shock waves. In [14, 18] Holian et al. pointed out the fact that even a 1D shock considered in a 3D system (a piston compression along a principal direction of a crystal for example) may not look like the typical 1D pattern of Figures 1 or 2. If the crystal is at zero temperature, then the compression pattern in 3D is the same as the 1D one, with a soliton train at the front. But if positive temperature effects are considered, the interactions of the particles with their neighbors - especially in the transverse directions - lead to the destruction of the coherent soliton train at the front, and a steady-regime can be reached (shock with constant thickness).

Therefore, 1D models are often supplemented with a postulated dissipation. The corresponding damping term in the equations of motion usually accounts for radiative damping [13, 24, 25], or may compensate thermal fluctuations [1] from an external heat bath for a system at equilibrium. Let us point out that purely dissipative models may stabilize shock fronts. However, temperature effects then completely disappear. In particular, no jump in kinetic temperature can be observed in purely dissipative 1D simulations. Besides, we also aim here at motivating the usually postulated dissipation and memory terms, and show that they arise naturally as effects of (conveniently chosen) higher dimensional degrees of freedom.

To the best of the author’s knowledge, there is no existing model that could both account for higher dimensional effects in non equilibrium dynamics and be mathematically tractable. We introduce a classical deterministic heat bath model, as an idealized way to couple the longitudinal modes of the atom chain to other modes. This model is justified to some extent by heuristic considerations in Section 3.1. We are then able to derive a generalized Langevin equation describing the evolution of the system, and recover a stochastic model in some limiting regime.
3.1. Form of the perturbations arising from higher dimensional degrees of freedom

Consider the system described in Figure 8, which is still a 1D atom chain, but where each particle in the 1D chain also interacts with two particles outside the horizontal line. These particles aim at mimicking some effects of transverse degrees of freedom. The transverse particles are placed in the middle of the springs and have only one degree of freedom, namely their ordinates $y_n$. The particles in the 1D chain are still assumed to have only one degree of freedom as well. This means that we constrain them to remain on the horizontal line. The interactions between the particles in the chain and the particles outside the chain are ruled by a pairwise interaction potential, for example the same potential as for interactions in the 1D chain.

Figure 8. Notations for the interaction of a transverse particle with particles on the 1D atom chain.

Consider small displacements around equilibrium positions. The pairwise interaction potentials can therefore be taken harmonic. Up to a normalization, and for a displacement $x$ from equilibrium position, $V(x) = \frac{1}{2}x^2$.

We first turn to the case $\theta = \frac{\pi}{3}$ corresponding to a 2D regular lattice. At first order,

$$d_n = \left[ \left( \frac{1}{2}(1 + x_{n+1} - x_n) \right)^2 + \left( \frac{\sqrt{3}}{2} + y_n \right)^2 \right]^{1/2} \approx 1 + \frac{1}{4}(x_{n+1} - x_n) + \frac{\sqrt{3}}{2}y_n.$$  

We now focus on the evolution of $x_n$. All the equalities written below have to be understood as equalities holding at first order in $O(|x_n|), O(|y_n|)$. Considering only interactions with the neighboring particles on the horizontal line, and the additional interaction with the particle $y_n$,

$$\ddot{x}_n = \frac{9}{8}(x_{n+1} - 2x_n + x_{n-1}) + \frac{\sqrt{3}}{4}(y_n - y_{n-1}).$$

The equation governing the evolution of $y_n$ is:

$$\ddot{y}_n = -\frac{3}{2}y_n - \frac{\sqrt{3}}{2}(x_{n+1} - x_n).$$
More generally, consider the system of Figure 8 with a general angle $\theta$. The equilibrium distance is now $x_0 = \frac{d_0}{2\cos \theta}$, and the corresponding normalized harmonic potential is $V(d) = \frac{1}{2}(\frac{d}{d_0} - 1)^2$.

The normalized distance $d_n = \frac{d_n}{d_0}$ is now

$$d_n = 1 + \cos^2 \theta (x_{n+1} - x_n) + 2 \sin \theta \cos \theta \cdot y_n.$$ 

The additional longitudinal force exerted on $x_n$ by $y_n$ is then

$$f_n = \cos^2 \theta [\cos \theta (x_{n+1} - x_n) + 2 \sin \theta \cdot y_n].$$

Summing over $N$ particles that do not interact with each other, each one being characterized by an angle $\theta_i$, the additional force on $x_n$ is seen to be of the form

$$F_n = A_N(x_{n+1} - 2x_n + x_{n-1}) + \sum_{i=1}^{N} K_i(y_n^i - y_{n-1}^i),$$

with $K_i = 2 \cos^2 \theta_i \sin \theta_i$ and $A_N = \sum_{i=1}^{N} \cos^3 \theta_i$. So, the equation of motion for $x_n$ is

$$\ddot{x}_n = (1 + A_N)(x_{n+1} - 2x_n + x_{n-1}) + \sum_{i=1}^{N} K_i(y_n^i - y_{n-1}^i).$$

(3.1)

The equations for the $y_n^i$ can be obtained in the same way as before:

$$\ddot{y}_n^i = -a_i y_n^i - 2K_i(x_{n+1} - x_n).$$

(3.2)

These linear perturbations are only valid for small displacements, i.e. when the approximation of the full potential by its harmonic part is justified. Notice moreover that we discard any type of interaction of the $y$ particles with each others.

However, this motivates an attempt to take into account missing degrees of freedom by introducing a heat bath whose form will lead to equation of motion similar to (3.1) - (3.2). We now turn to this task.

3.2. Description of the heat bath model

We consider the following Hamiltonian for a coupled system consisting of the system under study (S) and a heat bath (B) described by bath variables $\{y_n^j\}$ ($n \in \mathbb{Z}$, $j = 1, \ldots, N$). To use a heat bath is classical but was never done in the context of 1D chains to the author’s knowledge. The full Hamiltonian reads:

$$H(\{q_n, p_n, \tilde{q}_n^j, \tilde{p}_n^j\}) = H_S(\{q_n, p_n\}) + H_{SB}(\{q_n, p_n, \tilde{q}_n^j, \tilde{p}_n^j\}).$$

(3.3)

where $(q_n, p_n, \tilde{q}_n^j, \tilde{p}_n^j) = (x_n, \dot{x}_n, y_n^j, m_j \dot{y}_n^j)$, $H_S$ is given by (2.2), and

$$H_{SB}(\{q_n, p_n, \tilde{q}_n^j, \tilde{p}_n^j\}) = \sum_{n=-\infty}^{\infty} \sum_{j=1}^{N} \frac{1}{2m_j} (\tilde{p}_n^j)^2 + \frac{1}{2} k_j [\gamma_j (q_{n+1} - q_n) + \tilde{q}_n^j]^2.$$  

(3.4)

The interpretation is as follows. Each spring length $\delta x_n = x_{n+1} - x_n$ is thermostated by a heat bath $\{y_n^j\}$, in the spirit of [7, 28]. The parameter $k_j$ is the spring constant of the $j$-th oscillator, $m_j$ its mass, $\gamma_j$ weights the coupling between $\Delta x_n$ and $y_n^j$. Note that although more general cases can be considered [22, 20], the coupling is taken bilinear in
the variables, for it allows for an exact mathematical treatment. Indeed, a generalized Langevin equation (GLE) can be easily recovered (see [7, 28] for seminal examples). To the author’s knowledge, it is also the only case where the limit $N \to \infty$ can be rigorously justified.

Other physical motivations may be presented, such as the representation of extra variables in Fourier modes leading to a Hamiltonian similar to (3.3), see [2]. These extra degrees of freedom allow for some "transverse" radiation of the energy.

3.3. Derivation of the generalized Langevin equation

3.3.1. General procedure

Up to a rescaling of $y^n_j$, we may assume that all masses $m_j$ are 1. The only parameters left for the coupling are the coupling factors $\gamma_j$. Introducing the pulsations $\omega_j$ given by $\omega_j = k_j^{1/2}$, the equations of motion read:

$$\ddot{x}_n = g_N(x_{n+1} - x_n) - g_N(x_n - x_{n-1}) + \sum_{j=1}^{N} \gamma_j \omega_j^2 (y^n_j - y^n_{j-1}),$$

(3.5)

$$\ddot{y}^n_j = -\omega_j^2 \left[ y^n_j + \gamma_j (x_{n+1} - x_n) \right],$$

(3.6)

where

$$g_N(x) = V'(x) + \left( \sum_{j=1}^{N} \gamma_j^2 \omega_j^2 \right) x.$$  

(3.7)

Notice the structural similarities of (3.5) with (3.1) and of (3.6) with (3.2).

The procedure is classical [28]. The solutions $\{y^n_j\}$ of (3.6) are integrated and then inserted in (3.5) for $\{x_n\}$. The integrability of the system is clear (once initial conditions in velocities and displacements are set) when the force $g_N$ is globally Lipschitz. This is for example the case when the sum $\sum_{j=1}^{N} \gamma_j^2 \omega_j^2$ is finite, and when $V'$ is globally Lipschitz, which is indeed true for the Toda potential (2.5). For the Lennard-Jones potential (2.4) it remains true as long as the energy of the system is finite (since the potential diverges when $x \to -1$, the bound on the total energy implies $x > x_0 > -1$, and a bound on the Lipschitz constant can be given by $V'(x_0)$).

The computation gives:

$$y^n_j(t) = y^n_j(0) \cos(\omega_j t) + \frac{\dot{y}^n_j(0)}{\omega_j} \sin(\omega_j t) + \int_0^t \gamma_j \omega_j \sin(\omega_j s)(x_{n+1} - x_n)(t - s) \, ds.$$  

Integrating by parts and inserting in (3.5):

$$\ddot{x}_n(t) = V'(x_{n+1} - x_n) - V'(x_n - x_{n-1})$$

$$+ \int_0^t K_N(s)(\dot{x}_{n+1} - 2\dot{x}_n + \dot{x}_{n-1})(t - s) \, ds + \tau^n_n(t),$$

(3.8)

where

$$K_N(t) = \sum_{j=1}^{N} \gamma_j^2 \omega_j^2 \cos(\omega_j t),$$
and
\[ r_N^n(t) = \sum_{j=1}^{N} \left( y_j^n(0) - y_{n-1}^j(0) \right) \gamma_j \omega_j^2 \cos(\omega_j t) + \left( \dot{y}_n^j(0) - \dot{y}_{n-1}^j(0) \right) \gamma_j \omega_j^2 \frac{\sin(\omega_j t)}{\omega_j} \]
\[ + \gamma_j^2 k_j \cos(\omega_j t)(x_{n+1} - 2x_n + x_{n-1})(0). \]

Formally, (3.8) looks like a GLE, provided \( r_N^n \) is a random forcing term. The dissipation term involves a memory kernel \( K_N \) and an “inner” friction \( \dot{x}_{n+1} - 2\dot{x}_n + \dot{x}_{n-1} \). The derivation made here shows that the usually postulated dissipation and memory arise naturally as effects of higher dimensional degrees of freedom. The dissipation term, classical in elasticity theory and postulated by some studies [13, 25], is derived here, as memory effects, that were also considered in [25], since the corresponding model was that of a viscoelastic material.

So, we are left with a description of the system only in terms of \( \{x_n\} \). To further specify the terms, we have to describe the choice of the heat bath spectrum \( \{\omega_j\} \), the coupling constant \( \gamma_j \) and the initial conditions for the bath variables.

### 3.3.2. Choice of the constants

We choose the values [21]:
\[ \omega_j = \Omega \left( \frac{j}{N} \right)^k, \quad \gamma_j^2 \omega_j^2 = \lambda^2 f^2(\omega_j) (\Delta \omega)_j, \quad f^2(\omega) = \frac{2\alpha}{\pi} \frac{1}{\alpha^2 + \omega^2}, \quad (3.9) \]
where \( (\Delta \omega)_j = \omega_{j+1} - \omega_j, \ \alpha, \lambda > 0 \) and \( k > 0 \).

The function \( f^2 \) is defined this way for reasons that will be made clear in Section 3.4.

The heat bath spectrum \( \{\omega_j\} \) is more dense as \( N \) increases. The exponent \( k \) accounts for the repartition of the pulsations. More general choices could be made, involving randomly chosen pulsations [21]. However, we restrict ourselves to the case of deterministic pulsations.

We emphasize here once again that the constants chosen and the form of the coupling are not new. A similar choice is made in [21]. The novelty is in the application to a 1D chain, where independent heat baths are considered, each heat bath corresponding to a spring length.

We now motivate (3.9). Notice that an upper bound to the heat bath spectrum is imposed. This is related to the discreteness of the medium. Indeed, for a system at rest with particles distant from 1, the higher pulsation allowed is \( \pi \), corresponding to an oscillatory motion of spatial period 2. When particles come closer (for example if the mean distance between particles is \( a < 1 \)), the higher pulsation increases to the value \( \frac{\pi}{a} \) since the lowest spatial period is now \( 2a \). Taking then lower bound \( d_m \) for the minimal distance between neighboring particles, we get an upper bound for the spectrum, namely \( \Omega = \frac{\pi}{d_m} \).

The choice of the coupling constants between the system and the bath is an important issue. The only purpose of the heat bath in a 1D shock simulation is to mimic some effects of dimensionality, such as energy transfer to the transverse modes. This energy transfer can be quantified using (3.6). Indeed, the total energy transfer for a harmonic oscillator of pulsation \( \omega \) subjected to an external forcing \( \sigma \) is known [2]. More precisely, consider the following harmonic oscillator:
\[ \ddot{z} + \omega^2 z = h(t), \quad (3.10) \]
where $h$ is an external time-dependent forcing term. Then the total energy transferred by the external forcing to the system (from $t = -\infty$ to $t = +\infty$ for a system at rest at $t = -\infty$) is $\Delta E = \frac{1}{2} |\dot{h}(\omega)|^2$. The energy transfer to the heat bath occurs as described by (3.6). This gives a total energy transfer for a spring $x_{n+1} - x_n$ considered initially at rest:

$$\Delta E_n = \frac{1}{2} \sum_{j=1}^{N} \gamma_j^2 \omega_j^4 |\Delta x_n(\omega_j)|^2. \quad (3.11)$$

As a first approximation, a shock profile can be described as a self-similar jump: $\Delta x_n(t) = \delta H(n - c t n)$, where $\delta < 0$ is the jump amplitude, $c$ the shock speed, and $H$ is the Heaviside function. Then, $|\Delta x_n(\omega)| = \omega^{-1}$. The energy transfer (3.11) is therefore

$$\Delta E_n = \frac{\delta^2}{2} \sum_{j=1}^{N} \gamma_j^2 \omega_j^2.$$

With the spectrum (3.9), the condition $\Delta E_n \to C$ with $0 < C < \infty$ is satisfied:

$$\Delta E_n = \frac{\delta^2 \lambda^2}{2} \sum_{j=1}^{N} f^2(\omega_j)(\Delta \omega)_j \to \frac{\delta^2 \lambda^2}{2} \int_0^\Omega f^2 = \lambda^2 \delta^2 \sigma(\Omega).$$

The last expression is bounded since $f^2$ is integrable (We recall $\int_0^\infty f^2 = 1$). The function $\sigma$ is a $C^\infty$ function. Notice that the above convergence results from the convergence of the Riemann sum appearing on the left.

### 3.3.3. Choice of the initial conditions.

We consider initial conditions $\{y_n^i(0), \dot{y}_n^i(0)\}$ randomly drawn from a Gibbs distribution with inverse temperature $\beta_y$. This distribution is conditioned by the initial data $\{x_n, \dot{x}_n\}$. More precisely, set

$$y_n^i(0) = -\gamma_j(x_{n+1} - x_n)(0) + (\beta_y k_j)^{-1/2} \xi_j^n,$$

$$\dot{y}_n^i(0) = (\beta_y)^{-1/2} \eta_j^n; \quad (3.12)$$

where $\xi_j^n, \eta_j^n \sim \mathcal{N}(0,1)$ are independently and identically distributed (i.i.d.) random Gaussian variables. With these choices,

$$r_n^N(t) = \frac{1}{\sqrt{\beta_y}} \sum_{j=1}^{N} \omega_j \gamma_j \cos(\omega_j t) (\xi_j^n - \xi_{n-1}^j) + \omega_j \gamma_j \sin(\omega_j t) (\eta_j^n - \eta_{n-1}^j). \quad (3.14)$$

The probability space is induced by the mutually independent sequences of i.i.d. random variables $\xi_j^n, \eta_j^n$. Denote $D$ the linear operator acting on sequences $Z = \{z_n\}$ through $DZ = \{z_n - z_{n-1}\}$. So,

$$r_n^N(t) = \frac{\lambda}{\sqrt{\beta_y}} \sum_{j=1}^{N} f(\omega_j) \cos(\omega_j t) D\xi_j^n + f(\omega_j) \sin(\omega_j t) D\eta_j^n (\Delta \omega)_j^{1/2}. \quad (3.14)$$

For fixed $N$, the above expressions give

$$\mathbb{E}(r_n^N(t)(r_n^N(s))^T) = \frac{1}{\beta_y} K_N(t - s)DD^T \quad (3.15)$$
where \( r^N = (\ldots, r^N, \text{dots}) \) and the linear operator \( DD^T \) acts on sequences \( Z \) as \( DD^T z = \{z_{n+1} - 2z_n + z_{n-1}\} \). This relation is known as the fluctuation-dissipation relation, linking the random forcing term and the memory kernel. Notice that the noise term is correlated both in time and in space. The behavior of the system when \( N \to \infty \) is then an interesting issue, that can help us to get a better understanding of the phenomenas at play.

3.4. Limit when \( N \to \infty \)

3.4.1. Limit of the dissipation term
The memory kernel can be seen as a Riemann sum. The limit is then:

\[
K_N(t) = \lambda^2 \sum_{j=1}^{N} f^2(\omega_j) \cos(\omega_j t)(\Delta\omega)_j \to \lambda^2 \int_0^\Omega f^2(\omega) \cos(\omega t) \, dt = \lambda^2 K_\Omega(t)
\]  

(3.16)

when \( N \to \infty \), the convergence holding in \( L^1[0, T] \), \( T > 0 \).

The special choice (3.9) implies \( K_\Omega(t) \to e^{-\beta t} \) when \( \Omega \to \infty \) in \( L^\infty(\mathbb{R}_+) \). The memory kernel is then exponentially decreasing.

3.4.2. Limit of the fluctuation term
The limit \( N \to \infty \) gives the convergence of the noise term in a weak sense in \( C[0, T] \) (see the Appendix and [21]) toward a stochastic integral:

\[
r^N_n(t) \to \lambda r^\Omega_n(t) = \frac{\lambda}{\sqrt{\beta y}} \int_0^\Omega f(\omega) \cos(\omega t) DdW^{n,1}_\omega + f(\omega) \sin(\omega t) DdW^{n,2}_\omega \]  

(3.17)

where \( W^{n,1}_\omega, W^{n,2}_\omega (n \in \mathbb{Z}) \) are independent standard Brownian motions.

3.4.3. Limit of the equation
Formally, a stochastic integro-differential equation (SIDE) is obtained in the limit \( N \to \infty \):

\[
\ddot{x}_n(t) = V'(x_{n+1} - x_n) - V'(x_n - x_{n-1}) + \lambda^2 \int_0^t K_\Omega(s)(\ddot{x}_{n+1} - 2\ddot{x}_n + \ddot{x}_{n-1})(t - s) \, ds + \lambda r^\Omega_n(t),
\]  

(3.18)

(3.19)

with

\[
K_\Omega(t) = \int_0^\Omega f^2(\omega) \cos(\omega t) \, d\omega,
\]  

\[
r^\Omega_n(t) = \frac{1}{\sqrt{\beta y}} \int_0^\Omega f(\omega) \cos(\omega t) DdW^{n,1}_\omega + f(\omega) \sin(\omega t) DdW^{n,2}_\omega,
\]

and the fluctuation-dissipation relation

\[
\mathbb{E}(r^\Omega(t)(r^\Omega(s))^T) = \frac{1}{\beta y} K_\Omega(t - s) DD^T,
\]  

(3.20)

where \( r^\Omega = (\ldots, r^\Omega, \ldots) \). The way the solutions of (3.8) converge to the solutions of (3.18) can be made rigorous by a direct adaptation of the results of [21]: the convergence of \( x^N_n \) solution of (3.8) to \( x_n \) solution of (3.18) is weak in \( C^2[0, T] \) (in the sense of continuous random processes). We refer to the Appendix for some precisions on the proof.
The random process $r_n^{\Omega}$ are Ornstein-Uhlenbeck (OU) processes. In order to precise these OU processes, the SIDE (3.18) can be rewritten as a stochastic differential equation (SDE). This is done in the limiting case $\Omega \to \infty$ where a Markovian limit can be recovered when considering an additional variable [21]. Notice that when $\Omega \to \infty$, $K_h(t) \to K(t) = e^{-\alpha t}$. Denoting $Q = (\ldots, x_{n-1}, x_n, x_{n+1}, \ldots)$, $P = (\ldots, \dot{x}_{n-1}, \dot{x}_n, \dot{x}_{n+1}, \ldots)$, $V(Q) = \sum_{n=-\infty}^{\infty} V(x_{n+1} - x_n)$ and $R = (\ldots, R_{n-1}, R_n, R_{n+1}, \ldots)$, the previous SIDE (3.18) is equivalent to the following SDE:

$$
\begin{align*}
\frac{dQ}{dt} &= P \\
\frac{dP}{dt} &= (R - \nabla V(Q)) dt \\
\frac{dR}{dt} &= -(\alpha R + \lambda^2 DD^T P) dt + \lambda \sqrt{\frac{2\alpha}{\beta}} DdW,
\end{align*}
$$

where $W$ is a standard Brownian motion, and with initial conditions $r_n(0) \sim \lambda \beta^{-1/2} N(0, 1)$.

The limiting equation (3.17) shows the main effects of the heat-bath interaction: The pure 1D equation (2.3) is supplemented by two terms, one dissipation term with an exponentially decreasing memory, and a random forcing. Therefore the heat bath acts first as an energy trap, absorbing some of the energy of the shock when it passes. This energy is then given back to the system through the random forcing term to an amount precised by (3.20). This allows the equilibration of the downstream domain. This heuristic interpretation is confirmed by some numerical simulations of (3.8) in Section 4.

3.5. Generalization of the system-bath interaction

The Hamiltonian of the system can be written in an abstract form as

$$
H(x, y_N) = \frac{1}{2} |\dot{x}|^2 + F(x) + \frac{1}{2} |My_N|^2 + \frac{1}{2} |Ax - By_N|^2
$$

(3.22)

where $x = (\ldots, x_{n-1}, x_n, x_{n+1}, \ldots)$ and $y_N = (\ldots, y_{n-1}^1, y_{n-1}^N, y_1^1, \ldots, y_n^N, \ldots)$. The matrix $M$ is a mass matrix (operator), $A$ and $B$ are general operators, $F(x) = \sum_{n=-\infty}^{\infty} V(x_{n+1} - x_n)$.

In the previous example, $B$ was diagonal. But more generally, $B$ could be considered as tridiagonal: this could model the interaction of two neighboring heat baths linked to neighboring spring lengths.

4. Numerical results

The equations of motion (3.5), (3.6) are integrated numerically for a given $N$, using a classical velocity-Verlet scheme. The system is initialized with velocities and displacements generated from (3.12) and (3.13) in the $y$-coordinates, and from (2.6) in the $x$ coordinates. Note that the quantities $\frac{1}{\beta_x}$ and $\frac{1}{\beta_y}$ may differ. The system is then first left to evolve freely, so that the coupling between transverse and longitudinal directions starts.

Shock waves are generated using a piston in the same fashion as in Section 2.2.3, giving Figures 9 and 10. We then study relaxation waves (Figure 11).
The time-step $\Delta t$ is chosen to ensure a relative energy conservation of $10^{-3}$ in the
absence of external forcing. Typically, $\Delta t = 0.01$. The spectrum density parameter $k$ in
(3.9) is taken to be $k = 1$. Other choices lead to the same kind of simulation results.

Notice that, if $L$ represents the size of the 1D chain, the algorithmic complexity scales
as $O(LN)$. The computations were made on an usual desktop computer (Pentium 1.0 GHz),
and only took about a couple of hours for the most demanding ones.

4.1. Sustained shock waves

Figures 9 and 10 show the different patterns obtained in the case of a system coupled to a
heat bath. Notice that the upper bound to the spectrum, $\Omega$, is of order $\pi$ since the shock
is not too strong, and hence the medium is not too compressed. The parameter $\alpha$ is taken
less or equal to $\Omega$ so that $K_\Omega$ and $\sigma(\Omega)$ are sufficiently close from their limiting values.

The parameter $\lambda$ was varied in the range $[0, 5]$. If $\lambda$ is too small, the coupling is too
weak and the profiles look like the pure 1D ones (Note that we recover the purely 1D
model with Hamiltonian (2.2) when $\lambda = 0$). If $\lambda$ is too high, the forcing may be too strong,
leading to the collapse of two neighboring particles if the time step is not small enough. A
good choice of $\lambda$ involves a good rate of energy transfer to the transverse modes. For the
moment the choice of $\lambda$ is completely empirical. It would be desirable to estimate it from
full 3D simulations. This is precised to some extent in the next Section.

The results show that the introduction of transverse degrees of freedom has important
consequences on the pure 1D pattern. The soliton train at the front is destroyed, and the
shock thickness is constant along time, instead of growing in time as in the pure 1D case.

This is to the author’s knowledge the first result of this kind for a 1D chain. Thus a
steady regime can now be reached, and these simulations really seem to deserve the name
“shock waves”. In contrast to the pure 1D model results, these simulations have now the
same qualitative behavior as 3D simulations or experiments.

Figure 9. Relative displacement profiles for the system coupled to a heat bath (left), and
comparison with a thermalized shock (right). For the thermalized shock, the parameters
are $u_p = 0.3$, $b = 10$ and $\frac{1}{\sqrt{\beta_p}} = 0.01$. For the system coupled to a heat bath, the
additional parameters are $\frac{1}{\sqrt{\beta_p}} = 0.02$, $\alpha = 5$, $\Omega = 10$, $\lambda = 0.5$. The number of transverse
oscillators is $N = 25$. 
4.2. Rarefaction waves

As can be seen in Figure 11, a rarefaction wave develops and progressively weakens the shock (notice that the velocities decrease and that the relative displacement increase compared to Figures 9 and 10). This is indeed the expected physical behavior for a viscous fluid. This dissipation can be interpreted as energy transfer to the transverse modes.

Besides, no soliton train survives, contrarily to the pure 1D case, where the solitons are not destroyed and move on unperturbed. In the pure 1D case, there is no weakening of the initial wave, only dispersion. Once again, to our knowledge, this is the first time a 1D discrete model behaves as expected.
5. Conclusion

This study indicates a possible track to thermostate a 1D lattice in a deterministic way, without fixing the temperature as would require a Langevin thermostetting for instance. Indeed, when the shock passes, the temperature changes, and a Langevin simulation asks for an a priori knowledge of the temperature in the shocked region.

The interactions of the chain and the bath naturally lead to memory effects, and can be described by a memory kernel, at least in some limiting regime. Numerical experiments illustrate the success of this method. This model indeed qualitatively reproduces some important features of shock waves (sharpness of the shock front, existence of relaxation waves, equilibration after the shock has passed). This is in contrast with the classical pure 1D model.

However, this heat-bath thermalization is better suited for shocks that are not too strong. On the other hand, for strong shocks, nonlinear effects should play an important role in the energy transfer in the transverse modes, and a bilinear coupling such as (3.4) may not be a relevant modelling. In this case, a nonlinear coupling in the spirit of [20] should be more adapted.

An interesting issue is now to compare those reduced 1D profiles with profiles arising from full 3D simulations. Figure 12 compares the velocity of the center of mass of a slice of constant thickness of a Lennard-Jones solid (using a FCC structure), in reduced units [16], and the velocity of the corresponding representative particle (for the 1D model (3.3) ) as a function of the time. The agreement is reasonable, except at the shock front, where oscillations remain. A better agreement can however be obtained when using the SDE formulation (3.21). This agreement can be precised in a more quantitative way by parameter estimation techniques, as described in [20]. This work is in progress.

![Figure 12](image-url). Left: Longitudinal velocity profile for the center of mass of one slice of constant thickness as a function of time (in reduced units). Right: Velocity of the corresponding representative particle using the model (3.3) with a Lennard-Jones potential, with parameters $u_p = 0.3$, $\frac{1}{\sqrt{\alpha \omega^2}} = \frac{1}{\sqrt{2\alpha}} = 0.01$, $\alpha = 2$, $\Omega = 5$, $\lambda = 1$, $N = 200$. 


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Appendix

The proof of the convergence of the solutions of (3.8) to the solutions of (3.18) can be done as in [21], by a straightforward extension to the multi-dimensional case (in order to deal with convergence of sequences).

Denote $x_n^N$ the solution of (3.8) for a given number $N$ of transverse variables. We set

$$ \delta x_n^N = x_{n+1}^N - x_n^N. $$

The solution of (3.18) is noted $x_n$. We set $\lambda = 1$ to simplify notations. The extension to more general values of $\lambda$ is straightforward.

The space of real sequences in noted $\mathcal{H} = \mathbb{R}^N$, and is equipped with the usual $l^\infty$-norm. For a sequence $z = \{z_n\} \in \mathcal{H}$:

$$ |z|_{l^\infty} = \sup_{n \in \mathbb{Z}} |z_n|. $$

The space $\mathcal{H}$ endowed with this norm is then a separable complete metric space.

Consider the array of spring lengths

$$ Q_N = \begin{pmatrix} \vdots \\ \delta x_n^N \\ \vdots \end{pmatrix}, $$

and the array of random forcing terms

$$ G_N = \frac{1}{\beta y} \begin{pmatrix} \vdots \\ r_n^N \\ \vdots \end{pmatrix}. $$

We similarly define $Q$ and $G$ for the sequence $\{x_n\}$. (Note that these definitions are different from the definitions in Section 3.4.3 since we consider here the reduced relative displacements $\delta x_n$ instead of the particles’ positions $x_n$.)

Recall that the linear operator $D$, acting on sequences $z = \{z_n\} \in \mathcal{H}$, is defined by $Dz = \{Dz_n\} = \{z_n - z_{n-1}\}$. It follows $|DD^Tz|_{l^\infty} \leq 4|z|_{l^\infty}$.

Equation (3.8) can be rewritten as (recall $\lambda = 1$)

$$ \ddot{Q}_N = DD^TF(Q_N) + \int_0^t K_N(s)DD^T\dot{Q}_N(t-s) \, ds + DG_N(t). $$
Introducing $K_N(t) = \int_0^t K_N(s) \, ds$ and integrating the convolution term by parts, equation (3.8) becomes

$$\ddot{Q}_N - \left( DD^T F(Q_N) + \int_0^t K_N(s) DD^T \dot{Q}_N(t-s) \, ds \right) = DG_N(t) - DD^T \dot{Q}_N(0) K_N(t). \quad \text{(A.1)}$$

This equation can be rewritten under a fixed point form as

$$(Id + R_N) \dot{Q}_N(t) = F_N(t). \quad \text{(A.2)}$$

As $F$ is Lipschitz, $\|R_N\|$ is small for small $T$. An usual Picard argument gives the existence and uniqueness of $\dot{Q}_N \in C([0,T], \mathcal{H})$ solving (A.2) for $T$ small enough (see [12], Section 12, for an analogous proof). Standard results also give the continuity of $\dot{Q}_N$ on $K_N \in L^1[0,T]$ and $U_N = DG_N - DD^T Q_N(0) K_N \in C([0,T], \mathcal{H})$. The mapping $(K_N, U_N) \mapsto Q_N$ is then continuous from $L^1[0,T] \times C([0,T], \mathcal{H})$ to $C([0,T], \mathcal{H})$ with the corresponding norms.

The convergence of $K_N$ in $L^1[0,T]$ is straightforward, and implies the convergence of $K_N$ in $L^1[0,T]$.

The convergence of $U_N$ results from the convergence of $K_N \in L^1[0,T]$ and from the convergence of $G_N$ to $G$ (in a way to precise). We refer to [11], Section VI.4., Theorem 2. Considering the collection of continuous real-valued stochastic processes $G_N$ with values in $\mathcal{H}$ (which is a separable complete metric space), we have to show:

(i) The finite-dimensional distributions of $G_N$ weakly converge to those of $G$, which is a continuous process.

(ii) A tightness inequality of the form

$$\forall t, t + u \in [0,T], \quad \mathbb{E} \left[ |G_N(t+u) - G_N(t)|^2 \right] \leq C|u|. \quad \text{Then it follows} \quad G_N \Rightarrow G \quad \text{in} \quad C([0,T], \mathcal{H})\text{-weak.}$$

These two points are straightforward generalizations of the proof in [21] (in the case of non-random pulsations $\omega_j$) when extended to sequences with values in $\mathcal{H}$, giving the convergence $U_N \Rightarrow U$ in $C([0,T], \mathcal{H})$-weak.

The convergences of $K_N$ to $K$ in $L^1[0,T]$ and $U_N$ to $U$ in $C([0,T], \mathcal{H})$ in a weak sense then give the convergence of $\dot{Q}_N$ in $C([0,T], \mathcal{H})$ in a weak sense. Therefore, $Q_N \Rightarrow Q$ in $C^2([0,T], \mathcal{H})$-weak. This implies the convergence in a weak sense for all the components of $Q_N$ for $T$ small enough.

For general $t$, consider $e^{-\gamma t} Q_N$ for $\gamma$ large enough, and rescale appropriately the operators appearing in (A.2). The proof then follows the same lines.