Phason Dynamics in One-Dimensional Lattices

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In quasicrystals, the phason degree of freedom and the inherent anharmonic potentials lead to complex dynamics which cannot be described by the usual phonon modes of motion. We have constructed simple one-dimensional model systems, the dynamic Fibonacci chain (DFC) and approximants thereof. They allow us to study the dynamics of periodic and quasiperiodic structures with anharmonic double well interactions both by analytical calculations and by molecular dynamics simulations. We found soliton modes like breathers and kink solitons and were able to obtain closed analytical solutions for special cases, which are in good agreement with our simulations. Calculation of the specific heat shows an increase above the Dulong-Petit value, which is due to the anharmonicity of the potential and not caused by the phason degree of freedom.

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I. INTRODUCTION

Quasicrystals are aperiodic crystals with incommensurate spatial frequencies due to noncrystallographic symmetries. Their structure can be modelled by the projection of a planar, narrow stripe (acceptance stripe) of a higher-dimensional crystal onto physical space. As a consequence, the number of basis vectors in reciprocal space is higher than the dimension of physical space, which leads to the possibility of new symmetries not allowed in periodic crystals like e.g. five-fold or icosahedral symmetry.

The dynamics of quasicrystals is governed by a complicated potential energy landscape with more minima than atoms. Most of the time the atoms stay in their respective local minima. However, on a picosecond time scale, they can overcome the energy barrier and swap into a neighboring minimum. To stress the instantaneous nature of this process, it is called a flip. The occurrence of these flips is a characteristic feature of quasicrystals and follows from the construction method in higher-dimensional space: If the acceptance stripe is translated perpendicular to the physical space, then some lattice points leave the stripe while others enter it. The result in physical space are discrete jumps of atoms. In the continuum picture, such fluctuations of the stripe are identified with internal degrees of freedom, the so-called phason modes. Together with the conventional phonon modes, they make up the dynamics of quasicrystals.

Although phason modes have great influence on macroscopic physical properties of quasicrystals, e.g. diffusivity, elasticity, plasticity, fracture, and phase transitions, there is little knowledge about the precise atomic motion underlying the phason flips. Nevertheless, phason fluctuations have been observed via speckle patterns and the tails of Bragg peaks in the structure factor. Furthermore, direct observations have been made by resolving the structural rearrangements of large atom clusters with high-resolution transmission electron microscopy. On the theoretical side, there are so far only stochastic and hydrodynamic modes, but few atomistic approaches.

To study phason modes closer, we have recently introduced a simple one dimensional model system, the dynamic Fibonacci chain (DFC) and its approximants. It consists of a chain of particles with classical interaction potentials that allow for phason flips. Depending on the initial conditions, the chain can be either periodic or quasiperiodic. In that paper, we focussed on the influence of phason flips on the dynamic and static structure factors in reciprocal space, and we could show that they mainly caused a broadening of the phonon peaks. The reason is the fundamental difference of phason flips and phonon modes: Phonons are periodic and extended in time and space, which yields clear signals in reciprocal space. In contrast, phasons are stochastically distributed. They can be observed only indirectly in reciprocal space by their interactions with phonons. The interaction is both static and dynamic. It is static, because the propagation of phonons is hindered by the disorder introduced due to previous phason flips. It is dynamic, since during the flip of a particle, the effective interaction is highly anharmonic and the harmonic approximation for phonon modes is not applicable. Together, these effects decrease the lifetime of phonons and broaden the characteristic lines.

In this paper we want to study the particle motion in real space. Can phason flips propagate? Are there soliton modes which other anharmonic systems show? These systems, for example the well known Frenkel-Kontorova (FK) chain or its continuous analogon, the sine-Gordon (SG) system, basically show three modes of motion: a) Kinks, that are “topological solitons” and describe a propagating topological defect in the chain (described as “translatorische Eigenbewegung” by Seeger); b) breathers, which are localized oscillating nonlinear soliton modes (“oszillatorische Eigenbewegung”); c) for small amplitudes, one can linearize the equations of motion and obtains the well known phonon modes. Indeed, we will show that upon strong excita-
tions the dynamic Fibonacci chain displays soliton modes in the form of breathers and kinks.

Furthermore, we calculate the specific heat of the DFC and observe that it rises beyond the Dulong-Petit law at high temperatures, as observed by Edagawa et al. in icosahedral Al-Mn-Pd. In section II, we introduce the model system we used for our studies. In section III, short chains are studied both by analytical approaches and by molecular dynamics simulations. Section IV describes the soliton modes and of thermodynamic properties of the DFC. In section V, we present our studies of solitary modes of energy minima) depends on the separation $d_n$ of the centers of the underlying double well potentials.

II. MODEL SYSTEM

The most simple quasiperiodic system is the one-dimensional Fibonacci chain. It can be constructed by projecting a stripe of the two-dimensional square lattice onto a straight line. There are two distinct distances of nearest neighbours in this chain, a short one ($S$) and a long one ($L$). Their ratio equals the golden number $\tau = \frac{1}{2}(1 + \sqrt{5}) \approx 1.618$. As the Fibonacci chain allows for flips of the $LS$ pair to $SL$ and vice versa, we choose it as the structure model for studying phason flips in quasicrystals. Another structure model we look at is the LS-chain, which is the periodic sequence of $L$ and $S$. It is the simplest approximant of the Fibonacci chain with the possibility of flips.

The interaction between neighboring particles is given by a model potential that allows flips of atoms in $LS$ environments (two minima) and disallows flips in $LL$ environments (one minimum) and $SS$ environments. Note that the latter do not occur in the perfect Fibonacci chain, but might exist in our systems.

It turns out that the best approach is an interaction potential of the form

$$V_{\text{part}}(x) = ax^4 + bx^2,$$

where we usually use $a = 1$ and $b = -2$. In general, it is always possible to transform an arbitrary potential of the form (1) to one with $a = 1$ and $b \in \{-2, 0, 2\}$ by a coordinate change. $b < 0$ is a double well with an energy barrier $E_0 = \frac{b^2}{4a}$, and $b \geq 0$ has a single minimum.

The effective potential of particle $n$ is a superposition of the double well potentials which left and right neighbour exert and is plotted in figure 1. $A$ is the length of the double well “spring” and, for our standard choice $a = 1, b = -2$, equals $2\tau + 1$. $d_n = x_{n+1} - x_{n-1} - 2A$ is the separation of the centers of the double wells. If we choose the coordinate origin in the middle of the neighbour particles, the potential reads:

$$V_{\text{eff}}(x_n) = V_{\text{part}}\left(x_n - \frac{d_n}{2}\right) + V_{\text{part}}\left(x_n + \frac{d_n}{2}\right)$$

$$= 2 \left(ax_n^4 + bx_n^2 + \Delta\right)$$

with $\tilde{b} = b + \frac{3}{2}ad_n^2$ and $\Delta = a(d_n/2)^4 + b(d_n/2)^2$. For $b < 0$ (double well), there is a critical separation $d_{\text{crit}}$, where the sign of $\tilde{b}$ changes:

$$d_{\text{crit}} = \sqrt{\frac{2|b|}{3a}}$$

So, there is a bifurcation: for $|d_n| < d_{\text{crit}}$, we have a double well with minima at $x_{\pm}(d_n) = \pm \sqrt{\frac{|b|}{2a} - \frac{3}{4}d_n^2}$; for $|d_n| > d_{\text{crit}}$, there is only one minimum at $x = 0$.

The total potential of the chain is

$$V = \sum_n V_{\text{part}}(x_{n+1} - x_n - A)$$

For $x_{n+1} - x_{n-1} = 2A = 2(2\tau + 1)$ one obtains a double well in $LS$ environment with $\tilde{L} = 2\tau + 2$ and $S = 2\tau$, for $x_{n+1} - x_{n-1} = 2A \pm 2$ one obtains single wells in $LL$ and $SS$ environments.

In the two minima regime, the barrier height and distance of the minima are maximal, decreasing continuously when $d_n$ is approaching the critical distance.

III. SHORT CHAINS

In our attempts to learn about analytically accessible solutions of the equations of motion for chains underlying the above interaction potential we started with short chains. If these are constructed with periodic boundary conditions, they also describe periodic modes of motion in infinite chains.

Since the short “chain” of two particles can be described as a single particle in a double well potential (or to be more precise: a potential of the form $V(x) =$
FIG. 2: Effective potential $V_{\text{eff}}(u_1)$ of a particle in the chain of three particles of length LSL with the double-well interaction potential $V_{\text{part}}(x) = x^4 - 2x^2$ if one particle is not moving ($D = 1$, $a = 1$, $b = -2$, $u_2 = 0$). As in previous work, we use arbitrary units in this and the following figures.

FIG. 3: Trajectory $x_1(t)$ of the first particle of the chain of three particles of length LSL with the double-well interaction potential $V_{\text{part}}(x) = x^4 - 2x^2$ if one particle is not moving ($D = 1$, $a = 1$, $b = -2$, $u_2 = 0$, $M = 1$) at three different effective energies. While there is only one trajectory for high ($E_{\text{eff}} > 0$) and low ($E_{\text{eff}} < -\frac{3}{12}$) energies, there are two possible trajectories in the intermediate regime, because a particle can be situated in one of the two energy minima shown in figure 3. The trajectories of the simulated particles are not distinguishable from the analytical results.

The only approach to decouple these equations for the system $a = 1$, $b = -2$, $D = 1$ (LSL-Sequence which allows phason flips, because of the double well potential) was setting $u_2 = 0$, i.e. one particle does not move at all. The simplified equation of motion

$$\ddot{u}_1 := -\frac{12}{M} \left( au_1^3 + u_1 \left( \frac{b}{2} + \frac{1}{3}a (2u_2 + u_1 + D)^2 \right) \right)$$

(7a)

$$\ddot{u}_2 := -\frac{12}{M} \left( au_2^3 + u_2 \left( \frac{b}{2} + \frac{1}{3}a (2u_1 + u_2 - D)^2 \right) \right)$$

(7b)

One approach to handle the hamiltonian

$$H(x, p) = \sum_{i=1}^{3} \frac{p_i^2}{2M} + V_{\text{part}}(x_3 - x_2 - A) + V_{\text{part}}(x_1 + L_C - x_3 - A) + V_{\text{part}}(x_2 - x_1 - A)$$

(5)

(later, we will use particle mass $M = 1$) is the coordinate transformation

$$\begin{pmatrix}
R_1 \\
R_2
\end{pmatrix} = \begin{pmatrix}
\frac{1}{3} & \frac{1}{3} & \frac{1}{3} \\
-\frac{1}{2} & 1 & -\frac{1}{2}
\end{pmatrix} \begin{pmatrix}
x_1 \\
x_2 \\
x_3
\end{pmatrix} + \frac{L_C}{2} \begin{pmatrix}
0 \\
1 \\
0
\end{pmatrix}$$

(6)

where $R$ represents the coordinate of the center of mass. The transformation leads to a coupled system of nonlinear differential equations:

$$\ddot{u}_1 := -\frac{12}{M} \left( au_1^3 + u_1 \left( \frac{b}{2} + \frac{1}{3}a (2u_2 + u_1 + D)^2 \right) \right)$$

(8)

and

$$\ddot{u}_2 := -\frac{12}{M} \left( au_2^3 + u_2 \left( \frac{b}{2} + \frac{1}{3}a (2u_1 + u_2 - D)^2 \right) \right)$$

The potential is not symmetric any more but has the zeroes $\frac{1 \pm \sqrt{3}i}{2}$, local minima $(-1; -\frac{3}{4})$, $(\frac{1}{2}; -\frac{3}{12})$, and the local maximum $(0; 0)$. The equation $E_{\text{eff}} = \frac{1}{2}M \ddot{u}_1^2 + V_{\text{eff}}(u_1)$ finally leads to a trajectory

$$u_1 = r_2 + \frac{r_2 - r_1}{C \sin^2 \left( (t - t_0) \Omega \right) m} - 1$$

(10)

with the roots $r_i$ of $E_{\text{eff}} - V(u) = 0$:

$$\prod_{i=1}^{4} (u - r_i) = u^4 + \frac{2}{3} u^3 - u^2 - \frac{1}{4} E_{\text{eff}}$$

(11)

and

$$C = \frac{r_4 - r_1}{r_3 - r_2}$$

(12)

$$m = \frac{r_2 - r_3 r_4 - r_1}{r_1 - r_3 r_4 - r_2}$$

(13)

$$\Omega = \sqrt{\frac{2(r_1 - r_3)(r_4 - r_2)}{M}}.$$  

(14)
It is remarkable that the parameter $m$ of the Jacobi elliptic function $\text{sn}^{22\text{[23]}}$ is not only energy dependent; it has also very untypical values: From the analysis of the possible roots of equation (11) follows that the usual condition $0 \leq m \leq 1$ does not hold, but the admitted values for $m$ in the complex plane can have the following forms:

$$m_1 = \varphi$$  \hspace{1cm} (15a)
$$m_2 = \frac{1}{2} + 4\varphi$$  \hspace{1cm} (15b)
$$m_3 = e^{i\varphi}$$  \hspace{1cm} (15c)
$$m_4 = 1 - e^{i\varphi}$$  \hspace{1cm} (15d)

with energy dependent $\varphi \in \mathbb{R}$. So, $m$ is not necessarily a real number between 0 and 1, but can lie on the straight line of $m_2$ in the complex plane, on the real axis, and on the circles $m_3$ and $m_4$. It is worth noting, that the oscillation frequency $\Omega$ is energy dependent, which is due to the anharmonicity of the potential. Figure 2 shows typical trajectories for different energies. There is only one trajectory for high and low energies, but in the intermediate regime there are two trajectories for the same energy, one for the particle motion in the deep minimum of the effective potential (figure 2a) and one for the other minimum.

The observed behavior is typical for the dynamics of quasicrystals. At lower energies, the particles oscillate in their local minima. At intermediate energies, the particles can sit in different minima, but usually cannot overcome the barriers inbetween.

As a more general approach to solve Eq. (5) for $D = 0$ (the chain has the length $3\lambda$), we used Jacobi coordinates which are described in the classical solution\textsuperscript{22\textsuperscript{[23]}} of the three body problem\textsuperscript{22\textsuperscript{[23]}}. After transforming to Jacobi coordinates

$$\begin{pmatrix} R \\ x \end{pmatrix} = \begin{pmatrix} \frac{1}{2} \sqrt{2} & \frac{1}{2} \\ -\frac{1}{6} \sqrt{6} & \frac{1}{2} \sqrt{6} \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \end{pmatrix} - \frac{\sqrt{2}}{3} L_C \begin{pmatrix} 0 \\ 1 \end{pmatrix}$$  \hspace{1cm} (16)

and then to polar coordinates, $x = r \cos \varphi$, $y = r \sin \varphi$, we get a simpler Hamiltonian:

$$H = \frac{1}{2} M r^2 + \frac{1}{2} M r^2 \varphi^2 + \frac{3}{2} M r^2 + \frac{9}{2} a r^4 + 3 b r^2$$  \hspace{1cm} (17)

$R$ and $\varphi$ are cyclic variables. Thus, $\dot{R}$ and $L_\varphi := M r^2 \ddot{\varphi}$ are constant. This leads to equation

$$E = \frac{1}{2} M r^2 + \frac{L_\varphi^2}{2 M r^2} + \frac{9}{2} a r^4 + 3 b r^2$$  \hspace{1cm} (18)

where $E$ already contains the motion of the center of mass.

So for $D = 0$, the motion of three particles in one dimension can be treated as the motion of one particle with generalized angular momentum $L_\varphi$ in two dimensions.

For $a = 1$ and $b = -2$ the solution is

$$r^2 = r_3 + (r_2 - r_3) \text{sn}^2 \left( \Omega \cdot (t - t_0) \left| m \right. \right)$$  \hspace{1cm} (19)

with

$$\Omega = 3 \sqrt{\frac{r_3 - r_1}{M}}, \quad m = \frac{r_2 - r_3}{r_1 - r_3}$$  \hspace{1cm} (20)

$r_i$ are the solutions of $\frac{9}{2} x^3 - 3 x^3 - E x + \frac{L_\varphi^2}{2 M} = 0$:

$$\prod_{i=1}^{3} (x - r_i) = x^3 - \frac{4}{3} x^3 - \frac{2}{9} E x + \frac{L_\varphi^2}{9 M}$$  \hspace{1cm} (21)

For $L_\varphi \neq 0$, we get the angle

$$\varphi = \frac{L_\varphi}{M} \int_{r_3 + (r_2 - r_3) \text{sn}^2 \left( \Omega \cdot (t - t_0) \left| m \right. \right)} \frac{dt}{\sqrt{r_3 - r_1}}$$  \hspace{1cm} (22)

As there is no closed solution of this integral, only the case $L_\varphi = 0$ has been solved. In this case, $\varphi$ is constant and equation (19) simplifies to

$$r = \tilde{C}(E) \text{cn} \left( \tilde{\Omega}(E) (t - t_0) \left| \tilde{m}(E) \right. \right)$$  \hspace{1cm} (23)

with

$$\tilde{C}(E) = \sqrt{\frac{2}{3} + \frac{1}{3} \sqrt{4 + 2 E}}$$  \hspace{1cm} (24)
$$\tilde{\Omega}(E) = \sqrt{\frac{6}{M} \sqrt{4 + 2 E}}$$  \hspace{1cm} (25)
$$\tilde{m}(E) = \frac{1}{2} + \frac{1}{\sqrt{4 + 2 E}}$$  \hspace{1cm} (26)
$$cn$$ is another Jacobi elliptic function.

Here, the oscillation frequency is energy dependent, again. $$r(t)$$ resembles the motion of one particle in a double well potential. The constant angle $$\varphi$$ determines the ratios of $$r(t)$$ and the amplitudes $$x_i(t)$$ of the three particles. Figure 4 shows typical trajectories in this system. For $$E < 0$$ the $$cn$$ function evolves to a $$dn$$ function, because of $$\tilde{m} > 1$$.

In this section, we obtained results for special modes of motion in our systems showing interesting features like energy dependent oscillation frequencies and disconnected trajectories of same energy in phase space. Closed analytical solutions were found for a few special cases of particles oscillating in phase which also describe collective modes of motions in large chains.

IV. PERIODIC LS CHAIN IN REAL SPACE

The next step towards the Fibonacci chain is the periodic LS chain, a chain of particles with alternating distances $$L$$ and $$S$$ and total potential as in equation [1] with $$a = 1$$ and $$b = -2$$. The periodicity admits analytical calculations. Further simulations suggest that the results may be relevant for chains that contain defects like SS or LL sequences or for the Fibonacci chain.

A. Basic modes

We study this chain both by analytical approaches and by a numerical solution of the equations of motion with molecular dynamics simulations. In both cases periodic macro boundary conditions are applied.

Figure 5(a) shows some exemplary trajectories: After providing one particle with an initial velocity, we observe three solitary waves: One breather and two kink solitons propagating to both sides of the chain. These modes indeed show characteristics of the well-known modes of the sine-Gordon model. For example, the solitons can cross each other without noticeable interaction (figure 5(b)). Other initial conditions can lead to soliton-soliton interaction, which is not present in the standard sine-Gordon model. In the figure, the breather decays into two kink solitons.

Besides the kinks and breathers, naturally there are further modes: The rippled curves at large $$t$$ are due to phonons in the system.

B. Kinks

Figure 7(a) shows the typical trajectories in a chain with a propagating kink soliton. For this simulation, initial conditions (figure 6) were chosen such that at $$t = t_0$$ the particles of index $$n = 1, 2, \ldots$$ form an LS chain, as do the particles $$n = -1, -2, \ldots$$, but particles $$n = 1$$ and $$n = 0$$ are separated by $$L - \frac{\Delta}{2}$$, as are particles $$n = -1$$ and $$n = 0$$. The tunable parameters are the jump distance $$\Delta$$ and the initial velocity $$v_0(t_0) =: v_{\text{max}}(t_0)$$ of particle 0.

The kink propagates by flipping atoms: For particles in $$SL$$ environment (seen in direction of kink propagation; e.g. particle $$n = 0$$ in the figure) one can observe jumps by $$\Delta$$, for particles in LS environment (particle 1) jumps by $$\Delta + S - L$$. During this process LS environments become SL environments and vice versa. Of course, figure 6 shows only an idealized, approximate picture of the time development of a kink soliton. If $$t = t_0$$ is the moment of a snapshot of kink propagation with maximal particle velocity $$v_{\text{max}}$$, the jumping particle $$n = 0$$ is very close to the center of the two neighbouring particles $$n = 1, -1$$. These particles also have finite (but low) velocities. Because such deviations are not taken into account when initializing the system, the kinks typically need a short time to develop.
Because of the discrete lattice one expects that kinks radiate. In figure 7(b), the total energies $E_n(t)$ for particles $n$ at time $t$ are shown. One can observe a kink emitting phonons. The flipping particle loses energy and the kink decelerates until it has decayed completely into phonons. During that process, the maximal particle velocity decreases first approximately linearly in time, independent of the excitation parameters $v_{\text{max}}$, $\Delta$ at $t = 0$, and decays rapidly after reaching a critical velocity between 5 and 6, as figure 7 suggests. The time dependence of the kink propagation velocity $v_{\text{kink}}$ is qualitatively the same as $v_{\text{max}}$ because the two quantities are almost linearly related independent of the excitation parameters. So, in our system, there is no discontinuous variation of the kink’s deceleration as one expects for the Frenkel Kontorova chain, where two “deceleration regimes” exist.

Unfortunately, we could not find any analytical approximation describing the propagation of kinks. It is evident that kinks can only exist at high energy scales compared to the energy barrier of the potential (which is 1 in the partial interaction potential (1)).

Breathers are localized oscillatory modes. Numerical experiments (figure 9(a)) show that they are unstable. They can be destroyed by interaction with phonons or by tiny changes in the particle configuration.

We observed two types of breathers: One (type 2) basically consists of two neighbouring particles oscillating in opposite directions. The other (type 1) consists of a central oscillating particle and its oscillating neighbours. For the latter (type 1), our simulations suggest the fol-
Fig. 8: Development of the maximal particle velocity $v_{\text{max}}(t)$ of kink solitons. Independent of the excitation parameters ($\Delta$, $v_{\text{max}}(0)$), the particle velocity decays nearly linearly in time until a critical velocity (about 5) is reached.

Following analytical approximation:

$$
x_{-2}(t) = c_1 \quad \text{(27a)}
$$

$$
x_{-1}(t) = c_2 - X(t) \quad \text{(27b)}
$$

$$
x_0(t) = c_3 + 2X(t) \quad \text{(27c)}
$$

$$
x_{+1}(t) = c_4 - X(t) \quad \text{(27d)}
$$

$$
x_{+2}(t) = c_5 \quad \text{(27e)}
$$

Conservation of energy and $V(x) = ax^4 + bx^2$, $c_3 = 0$, $c_4 = A$, $c_5 = 2A$, $c_2 = -A$, and $c_1 = -2A$ (this is due to numerical observations) lead to

$$
E = 3M \dot{X}(t)^2 + 164aX(t)^4 + 20bX(t)^2 \quad \text{(28)}
$$

The solution for the standard double well potential ($a = 1$ and $b = -2$) is

$$
X(t) = \pm \sqrt{\frac{E}{3M}} \Omega^{-1} \text{sd}(\Omega(t - t_0)|m) \quad \text{(29)}
$$

with

$$
\Omega = \sqrt{\frac{40}{3M}} \sqrt{1 + \frac{41}{100}E} \quad \text{(30)}
$$

$$
m = \frac{1}{2} \left(1 + \frac{1}{\sqrt{1 + \frac{41}{100}E}}\right) \quad \text{(31)}
$$

The other type of breather (type 2) can be described by

$$
x_{-\frac{1}{2}}(t) = -\frac{3}{2}A \quad \text{(32a)}
$$

$$
x_{-\frac{1}{2}}(t) = -\frac{1}{2}A - X(t) \quad \text{(32b)}
$$

$$
x_{+\frac{1}{2}}(t) = +\frac{3}{2}A \quad \text{(32d)}
$$

which has this solution for our potential:

$$
X(t) = \pm \sqrt{\frac{E}{M}} \Omega^{-1} \text{sd}(\Omega(t - t_0)|m) \quad \text{(33)}
$$

Fig. 9: Simulated Breathers: Energy

(a) A rapidly decaying breather (type 1). The breather was constructed by combining a periodic LS chain, five particles of distance $A$ with initial velocities $0, −3.5114, 7.0228, −3.5114, 0$, and another periodic LS chain.

(b) Under ideal conditions (damped phonons, exact particle initialization) there exist long living breathers. This curve shows the energy profile of a breather of type 2 at $t = 1.5 \cdot 10^6$. 

with
\[ \Omega = \sqrt{\frac{12}{M} \sqrt{1 + \frac{1}{2}E}} \]  
\[ m = \frac{1}{2} \left( 1 + \frac{1}{\sqrt{1 + \frac{1}{2}E}} \right) \]

Although this localized ansatz is no stable solution, numerical simulations show good agreement for short times (figure 10). If the system is set up carefully and phonons, which would destroy the breather, are damped out, the system stabilizes when the neighbouring atoms oscillate with very low amplitudes. Figure 9(b) e.g. shows the energy profile in the chain at \( t = 1.5 \times 10^{-4} \). The high value of \( t \) demonstrates how stable breathers can be under special conditions.

V. DYNAMIC FIBONACCI CHAIN

In this section we present simulations of the dynamic Fibonacci chain, i.e. a chain of atoms whose initial positions are that of particles in the Fibonacci chain using our usual interaction potential with \( a = 1 \) and \( b = -2 \).

We set the initial velocities to random numbers that are normally distributed and allow us to simulate finite temperatures.

We prefer a temperature regulation by setting \( \langle E \rangle \) to a defined value and waiting for the system to reach an equilibrium state, since numerical thermostats can change the behaviour of the system and because there is no simple relation between \( T \) and \( \langle E \rangle \).

A. Ground state

If the length of the chain is a sum of integer multiples of \( L \) and \( S \), then all particles can sit in potential minima, which is a ground state of the system. For a fixed chain length or fixed periodic boundary conditions, the total number of \( L \) and \( S \) remains fixed because of their irrational ratio. All rearrangements by flips (\( LS \) to \( SL \) and vice versa) lead to energetically equivalent states.

The Fibonacci chain is a special case defined by \( p(L) = \tau p(S) \), where \( p(L) \) and \( p(S) \) are the frequencies of occurrence of \( L \) and \( S \) segments in the chain. Furthermore, the Fibonacci chain is the most uniform distribution of these two tile types within the class of all chains with the ratio \( \tau \). At finite temperatures, the chain will become disordered by flips, and defects like \( SS \) will form that are not present in the original Fibonacci chain. In thermodynamical equilibrium, the Fibonacci chain will approach a random tiling.

For low energies, the particles basically feel the parabolic potential that results from the Taylor expansion of the potential around its minima (i.e. \( \langle E_{\text{kin}} \rangle \approx \frac{1}{2} \langle E \rangle \)). The quartic term only dominates in the high energy regime \( E > 1 \) (\( \langle E_{\text{kin}} \rangle \approx \frac{3}{2} \langle E \rangle \)). As we will see, the nonparabolicity leads to significant deviations of the specific heat from Dulong-petit law.

In the following two sections, we will study the randomization of the Fibonacci chain and its heat capacity.

B. Randomization

While most observables we studied in the system showed quite plausible behaviour throughout our simulations (e.g. the flip frequency discussed below), we observed strange transitions of behaviour for such observables that depend on the number of \( LL, LS \) and \( SS \) environments in the system. The reason are structural changes in the chain that can be observed for high temperatures or long simulation times, when the Fibonacci chain has evolved to a random tiling.

The Fibonacci chain has another ratio of the possible environments
\[ p(SS) : p(LS) : p(LL) = 0 : \tau^{-1} : \tau^{-2} \]  
with
\[ p(L) : p(S) = \tau^{-1} : \tau^{-2} \]  

than the random tiling
\[ p(SS) : p(LS) : p(LL) = \tau^{-4} : 2\tau^{-3} : \tau^{-2} \]

Therefore observables such as the average frequency of particle transitions through the potential centers (zero for \( LS \) environments because the particles are trapped in one half of the double wells) change to an equilibrium state.
value until the random tiling is reached, following an exponential law

\[ f(t) = f_\infty + (f_0 - f_\infty)e^{-t/\tau_r} \]  \hspace{1cm} (39)

with the “randomization time” \( \tau_r \), see figure 11(a).

Figure 11(b) shows, that this randomization time follows an Arrhenius law

\[ \tau_r(T)^{-1} \propto e^{-E_r/kT} \] \hspace{1cm} (40)

where we find the “relaxation energy” \( E_r \approx 0.95 \).

C. Specific heat

Because of the degenerate ground states, flips do not contribute to entropy and hence, to the specific heat, which is solely determined by the interaction potential. We will prove this statement in the following.

Figure 12 shows for the ratio \( \langle E_{\text{pot}} \rangle / \langle E \rangle \) that thermodynamical properties of the dynamic Fibonacci chain agree remarkably with those of particles in a double well potential. It also demonstrates the transition from harmonic behaviour \( \langle E_{\text{pot}} \rangle \approx \frac{1}{2} \langle E \rangle \) to quartic behaviour \( \langle E_{\text{pot}} \rangle \approx \frac{1}{4} \langle E \rangle \) very well. In the relevant temperature range \( 0.1 \lesssim T \lesssim 1 \) for our simulations of the Fibonacci chain we are close to parabolic behaviour but already expect effects of anharmonicity.

Therefore, we use the analytically accessible system of noninteracting particles in a double well potential to determine the specific heat: The contribution of the po-

FIG. 12: The ratio of average potential energy and average total energy in the chain is the same as for non interacting particles in a double well potential. Thermodynamic properties are governed by the anharmonic potential.

FIG. 13: Temperature dependence of the specific heat of non interacting particles in a double well potential obtained by analytical calculation.

FIG. 11: Relaxation of the dynamic Fibonacci chain
tential energy can be calculated from

\[ C_{V,\text{pot}} = \frac{\partial \langle E_{\text{pot}} \rangle}{\partial T} = \frac{1}{kT^2} \left( \langle E_{\text{pot}}^2 \rangle - \langle E_{\text{pot}} \rangle^2 \right) \]  

Using

\[ \langle E_{\text{pot}} \rangle = E_0 \frac{A_2^- (\vartheta)}{A_0 (\vartheta)} \]  
\[ \langle E_{\text{pot}}^2 \rangle = E_0^2 \frac{A_4^- (\vartheta) - 4A_3^- (\vartheta) + 4A_2^- (\vartheta) + 2}{A_0 (\vartheta)} \]

with \( \vartheta := kT/E_0 \) and the integrals

\[ A_n^\pm (a) := \int_{-\infty}^{\infty} x^{2n} e^{-x^2/2a^2} \, dx \]

\[ = \left( \frac{a}{2} \right)^{2n+1} \Gamma \left( n + \frac{1}{2} \right) e^{\pm U} \left( n, \pm \sqrt{\frac{1}{2a}} \right) \]

where \( U \) is a parabolic cylinder function\(^{23}\). The plot of this function (figure 13) shows a rather interesting transition between harmonic and quartic regime: With increasing temperature \( C_{V,\text{pot}} \) rises from the parabolic \( 1/2k \) before approaching the quartic limit \( 1/k \), which has already been discussed in literature\(^{23}\). As the contribution of the kinetic energy \( C_{V,\text{kin}} = 1/k \) is constant, the total specific heat rises from the parabolic value \( k \) before declining and finally approaching quartic \( 1/k \) from beneath.

Edagawa et al\(^{20,29}\) observed that for high temperatures, \( C_V \) of Al\(_{63}\)Cu\(_{25}\)Fe\(_{12}\) rises above 3\( k \) which one expects from Dulong Petit. It is interesting that even for our simple model an increase of the specific heat by 14% can be observed, which is only caused by the interaction potential. This observation is in good agreement with the results of Grabowski et al\(^{20}\) which suggest that anharmonic interaction potentials cause high values of specific heat even for elementary fcc metals as Aluminium.

As seen above, no contribution of phasons to the specific heat is observed at high temperatures. Wälti et al. have measured an excess specific heat at low temperatures and assume its origin to lie in nonpropagating lattice excitations\(^{21}\). In our system we expect no influence of phasons on the low temperature specific heat, too, due to the low energy cutoff for the phason flips (Fig. 15). Low energy nonacoustic phonons are not present in the DFC, but in the asymmetric Fibonacci chain (AFC, Engel et al\(^{17}\)), which we are not dealing with here. The AFC is characterized by a bias in the double well which causes anticrossings in the many branches of the dynamical structure factor and leads to low lying flat bands (Fig. 15 of Engel et al\(^{17}\)).

To understand flips better, we calculated the energy distribution of particles in the moment a flip occurred.

There are two types of flip processes: “Autonomous flips” when atoms have enough energy to cross the potential barrier (figure 14(a)). For “induced flips” on the other hand, the flipping particle does not have to move at all: Particle motions of the next neighbours change the local energy landscape and cause the energy barrier to disappear. When a new barrier rises at the other side of the particle, the particle has completed a flip (figure 14(b)).

We calculated the average energies of particles in the moment of autonomous flips or of induced flips (here the “moment of the flip” is somewhat arbitrary, one can choose any point in the period of time when no barrier exists; we chose the moment when the energy barrier reappears). The particle energies were calculated as sum of kinetic energy and the local potential energy equation (2) shifted by \( 2 \cdot \frac{E_{\text{crit}}}{a} \), which causes the minimal energy of the double wells to be zero.

The probability distributions (figures 15(a) and 15(b)) display two prominent features: There is a sharp peak at \( E = 2 \), which is exactly the energy of the potential barrier. It is explained by the vanishing slope of \( V(x) \) there and the corresponding high density of states.

The other interesting point is the sharp cutoff at \( E \approx 0.9 \) which exists for both autonomous and induced flips. This lower threshold energy can be explained as the minimal potential energy of the energy barrier, i.e. the potential energy at the potential center in the case where the neighbouring atoms just reach the critical distance (equation (3)). This “minimal flip energy” is calculated to \( E_{\text{crit}} = \frac{1}{2} \).

It is remarkable that this minimal flip energy is close to the relaxation energy of equation (40).
The particle energies of atoms in the moment of a flip show two interesting features: A lower threshold energy $E \approx 0.9$ and a peak at $E \approx 2$.

The energy distribution for autonomous flips shows the threshold energy and the peak even better because time and energy of induced flips are somewhat difficult to define.

When studying energy transport in the chain at thermodynamical equilibrium and at low energies (i.e. energies lower than the potential barriers), we could not observe any prominent soliton modes in accordance with the results of section [IV] on the $LS$ chain.

On the other hand we could find low energy modes which can be described as regions of high energies propagating with constant velocity (they appear as straight lines in figure [16]). Because there is a lower threshold energy for flips, one expects that flips occur essentially along these propagating energy packets.

To verify this assumption, we marked the flips in the figure: Autonomous flips are represented by crosses (+), induced flips by circles ($\circ$). As it turns out, flips are indeed concentrated close to these modes.

When we increase temperature (i.e. $\langle E \rangle$), more and more of these high energy regions cross the energy threshold and cause a rapidly growing number of flips. Consequently, the flip frequency (number of flips per unit time and particle) is expected to follow an Arrhenius law.

Figure [17] shows the Arrhenius plot of the rapidly increasing frequencies we measured in our simulated system. As the straight lines fit our data nicely, the plot confirms our assumption. The slopes suggest activation energies 0.96, 1.05, 0.98 for induced, autonomous, and all flips which are fairly close to the minimal flip energy.
F. Solitons in the dynamic Fibonacci Chain

Next, we investigate the response of the chain to local excitations. All particles start on their equilibrium positions with zero velocity. Then, at time $t_0$, one particle $n_0$ is ‘kicked’ by setting $v_n(t_0) = v_0 \delta_{n,n_0}$.

By varying the excitation strength and excitation shape function, we can generate single solitons and breathers. As seen in Fig. 18(a), solitons usually pass each other with only little interaction. However it is also found that solitons are not stable. They continuously radiate phonons, therefore losing energy during propagation and slowing down. A soliton cannot exist with energy under a certain minimum threshold and will eventually decay. Similar results hold for breathers. In Fig. 18(b) a stationary one is shown. After radiating phonons for some time, it disappears at $t = 80$.

Note that the mathematical notion of soliton and breather is reserved for localized modes that do not decay in time. So, strictly speaking our modes are not ideal solitons/breathers. However, compared to the characteristic time of the system, both are stable and thus we adopt the naming. Especially in the case of high amplitude modes or in periodic chains (e.g. the periodic $LS$-chain), solitons and breathers are stable over very long times.

VI. CONCLUSION

The aim of our studies was an improved understanding of phasonic flips on an atomistic level and of the consequences of the anharmonic potentials that are typical for quasicrystals and other systems which provide a complex energy landscape.

Our main model system, the dynamic Fibonacci chain\cite{18} is constructed by providing particles assembled as a part of the one-dimensional quasiperiodic Fibonacci chain with an anharmonic pair potential of the form $V(x) = x^4 - 2x^2$ (double well potential). This system and simplified systems derived from it (short chains, periodic $LS$ chain) were studied both analytically and numerically using molecular dynamics simulations.

For the short “chains” consisting of three particles we found analytical solutions of the equations of motion with all particles oscillating in phase which can also be used to describe collective modes of motion in longer chains. As one might expect, the oscillation frequencies of these modes are energy dependent because of the anharmonic potential.

For studying the basic modes of motion which this potential induces we used the periodic $LS$ chain at $T = 0$. It turned out that the system does not only show the usual phonons but also two different soliton modes in an energy regime higher than the energy barrier of the potential: Breathers which are a localized oscillatory mode and kink solitons which can be viewed as a propagating topological defect ($LS$ environments flip to $SL$ environments and vice versa), or a propagating phason flip.

These modes are quite unstable and decay quickly. The kink solitons lose energy while propagating through the discrete lattice. Breathers also radiate when propagating through the chain, but tend to get trapped at a certain particle position; they can be kept stable in a very defined and shielded environment.

We did not find any analytical description of the kink solitons, but we succeeded in finding an analytical approximate solution describing a breather.

Our studies of the dynamic Fibonacci chain in thermodynamical equilibrium showed that there exists a lower threshold energy $\frac{8}{3}$ for atomic flips (for both flip types: autonomous flips and induced flips) which determines many properties of the system. E.g. it is important for transition of the Fibonacci chain to a random tiling (our potential does not penalize $SS$ environments which are not allowed in the Fibonacci chain!). We also found localized modes, which allow energy to be transported through the chain. If the temperature gets higher, an increasing number of these modes reaches the threshold energy, which leads to a large number of flips: The flip frequencies follow an Arrhenius law for low temperatures.

In the dynamic Fibonacci chain, we observed the same modes of motion as in the $LS$ chain. For observing soli-
tary modes such as breathers and kinks, the interaction potential is more important than the initial configuration. The same holds for thermodynamic properties: Specific heat increases above the harmonic value solely because of the interaction potential. Thus it is not the phason degree of freedom but the nonlinearity of the potential which causes the rise above the Dulong-Petit value.

The standard hydrodynamic theory of quasicrystals predicts only diffusive phasons. If one is leaving this long wavelength harmonic theory and proceeds to high excitation, then propagating modes appear in our model. It is an interesting question, whether such modes will be observable in highly excited quasicrystals.

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