

Surmounting collectively oscillating bottlenecks

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Ever since the seminal work by Kramers (for a comprehensive review see ref. [1]) we witness a continual interest in the dynamics of escape processes of single particles, of coupled degrees of freedom or of chains of coupled objects out of metastable states. To accomplish the escape the considered objects must cross an energetic barrier, separating the local potential minimum from a neighboring attracting domain. From the perspective of statistical physics mainly the thermally activated escape, based on the permanent interaction of the considered system with a heat bath, has been studied [1]. The coupling to the heat bath causes dissipation and local energy fluctuations and the escape process is conditioned on the creation of a rare, optimal fluctuation which in turn triggers an escape. To put it differently, an optimal fluctuation transfers sufficient energy to the system so that the system is able to statistically surmount the energetic bottleneck associated with the transition state. Characteristic time-scales of such a process are determined by the inverse of corresponding rates of escape out of the domain of attraction. Within this topic, numerous extensions of Kramers escape theory and of first passage time problems have been widely investigated [1,2]. Early

generalizations to multi-dimensional systems date back to the late 1960s [3]. This method is by now well established and is commonly put to use in biophysical contexts and for great many other applications occurring in physics and chemistry and related areas [4–14].

In order that the system comprised of coupled units may pass through a transition state an activation energy E_{act} has to be concentrated in the corresponding critical localized mode (LM). In view of controlling the process of barrier crossing we intend to demonstrate that the formation of the critical LM can be distinctly accelerated via the application of a weak external ac-driving. By use of optimally oscillating barrier configurations it is feasible that a far faster escape can be promoted, leading to a drastic enhancement of the escape dynamics. Particularly at low temperatures, where the rate of thermal barrier crossing is exponentially suppressed, such a scenario can be very beneficial.

Prior studies mainly dealt with the appearance of LMs in damped, driven deterministic nonlinear lattice systems [15–19]. Furthermore, the spontaneous formation of LMs (breathers) from thermal fluctuations in lattice systems, when thermalized with the Nosé method [20] has been demonstrated in [21,22]. Here we explain LM formation in a stochastic system involving dissipation in

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the presence of enduring spatio-temporal random forcing. In addition a weak external ac-field is applied rendering coherently *oscillating barriers*. The energy is introduced in the lattice coherently in the form of a plane-wave excitation as the response to the external ac-field and non-coherently through thermal fluctuations. We shall demonstrate that the stochastic source and the external ac-field conspire to produce such an instability mechanism of the stationary flat-state (plane-wave) solution yielding a spatially localized system state. Most importantly the formed LMs prove to be robust despite the continuously impacting thermal forces.

It should be noted that thermally activated escape of ensembles of non-interacting (individual) particles over a metastable potential landscape that is additionally subjected to either stochastic or coherent perturbations in the form of fluctuations or periodic driving has been studied in the prior literature, *e.g.* see in refs. [23–25]. For a comprehensive overview we refer the reader to ref. [26]. In particular, a resonant activation is observed, *i.e.*, the mean escape time (or the rate of escape [27]) attains a minimum (maximum) as a function of the correlation time of the fluctuations or the temporal driving period of the underlying potential variations. Moreover, the kink drift motion induced by oscillating external fields needs to be mentioned in this context [28]. Concerning a system of coupled elements the kink-antikink nucleation within a ϕ^4 chain model subjected to a deterministic periodic signal and uncorrelated noise has been studied in [5]. For optimal noise and coupling strength spatiotemporal (array enhanced) stochastic resonance is observed in the array of overdamped coupled elements. With the present study we focus on the collective nature of the ac-driven escape process of interacting weakly damped particles.

In detail, we study a one-dimensional lattice of damped nonlinear and ac-driven coupled oscillators which are subjected additionally to a heat bath at temperature T . Throughout the following we shall work with dimensionless parameters, as obtained after appropriate scaling of the corresponding physical quantities. The coordinate q of each individual nonlinear oscillator with a unit mass evolves in a cubic, single well on-site potential of the form

$$U(q) = \frac{\omega_0^2}{2}q^2 - \frac{a}{3}q^3. \quad (1)$$

This potential possesses a metastable equilibrium at $q_{min} = 0$, corresponding to the rest energy $E_{min} = 0$ and exhibits a maximum that is located at $q_{max} = \omega_0^2/a$ with energy $E_{max} \equiv \Delta E = \omega_0^6/(6a^2)$. Thus, in order for particles to escape from the potential well of depth ΔE over the energy barrier and subsequently into the range $q > q_{max}$, a sufficient amount of energy need to be supplied. The lattice dynamics is governed by the following system of coupled Langevin equations:

$$\ddot{q}_n + \gamma \dot{q}_n + \omega_0^2 q_n - a q_n^2 + \xi_n(t) - \kappa [q_{n+1} + q_{n-1} - 2q_n] - f \sin(\omega t + \theta_0) = 0. \quad (2)$$

The coordinates $q_n(t)$ quantify the displacement of the oscillator in the local on-site potential U at lattice site $n \in [1, N]$. The oscillators, referred to as “units”, are coupled bi-linearly to their neighbors with interaction strength κ . The friction strength is measured by the parameter γ and $\xi_n(t)$ denotes a Gaussian distributed thermal, white noise of vanishing mean $\langle \xi_n(t) \rangle = 0$, obeying the well-known fluctuation-dissipation relation

$$\langle \xi_n(t) \xi_{n'}(t') \rangle = 2\gamma k_B T \delta_{n,n'} \delta(t - t'), \quad (3)$$

with k_B denoting the Boltzmann constant. A homogeneous external periodic modulation field of amplitude f , frequency ω and phase θ_0 globally acts upon the system. In this work we use periodic boundary conditions according to $q_{N+1} = q_1$ and fix the parameters of the potential as follows: $\omega_0^2 = 2$ and $a = 1$, yielding $\Delta E = 4/3$. A deterministic escape scenario in the conservative, undriven limit of system (2) has been explored by us in [29,30].

To analyze the nonlinear character of the solutions of eq. (2) we first discard the noise ($\xi_n = 0$) and derive a nonlinear damped and driven discrete Schrödinger equation for the slowly varying envelope solution, $u_n(t)$, following the reasoning in [31], *i.e.*,

$$2i\omega_0 \dot{u}_n + i\gamma\omega_0 u_n + \kappa [u_{n+1} + u_{n-1} - 2u_n] + \alpha |u_n|^2 u_n + \frac{1}{2} f \exp[-i\Delta\omega t + \theta_0] = 0, \quad (4)$$

with the nonlinearity parameter reading $\alpha = 10a^2/(3\omega_0^2)$ and $\Delta\omega = \omega - \omega_0$. For the amplitude u_0 of a spatially homogeneous solution of eq. (4) of the form

$$u_n(t) = u_0 \exp[-i(\Delta\omega t + \theta_0)] + \text{c.c.}, \quad (5)$$

one obtains

$$\left[(2\omega_0 \Delta\omega + \alpha u_0^2)^2 + \gamma^2 \omega_0^2 \right] u_0^2 = \frac{1}{4} f^2. \quad (6)$$

In fig. 1 we depict the amplitude u_0 of the response versus the driving frequency curve for two different values of the driving amplitude f . At a bifurcation point a “jump” resonance related with a saddle-node bifurcation occurs and in certain range of the driving frequency multistability exists. In comparison for the larger driving amplitude, $f = 0.2$, the bifurcation point for the “jump” resonance occurs at a lower frequency value than for the driving with $f = 0.15$. Moreover, in the former case the system responds overall with higher amplitudes u_0 than in the latter. Notice that the system responds with large amplitude only within a frequency window and large amplitudes are obtained for a driving frequency lying below the band of linear frequencies, *viz.* $\omega < \omega_0 = \sqrt{2} = 1.414 \dots$ Similarly, for the response of the amplitude with regard to the driving strength f multistable solutions are possible as depicted with the inset in fig. 1. In order to investigate the stability of the homogeneous solution of eq. (2) we use

$$q_n(t; k = 0) = x(t) = u_0 e^{-i(\omega t + \theta_0)} + \frac{a}{\omega_0^2} \left[2 - \frac{1}{3} e^{-2i(\omega t + \theta_0)} \right] u_0^2 + \text{c.c.} \quad (7)$$

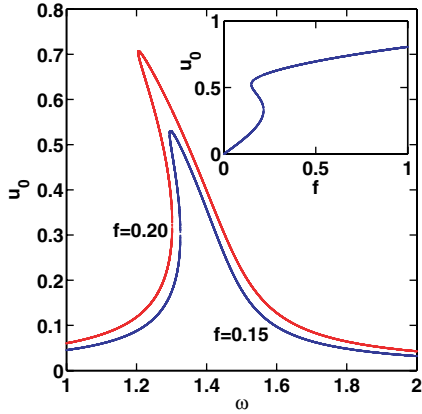


Fig. 1: (Color online) Response of the amplitude u_0 with respect to the frequency ω of the driving force for two fixed values of the driving amplitude as indicated in the plot. The damping constant is $\gamma=0.1$. The inset shows the response of the amplitude u_0 with respect to the driving amplitude f for a fixed frequency $\omega=1.295$. The remaining parameter values are $\omega_0^2=2$ and $a=1$.

and write with respect to the spatial perturbations A_n : $q_n(t) = x(t) + A_n(t)$. Since we impose periodic boundary conditions the Fourier-series expansion $A_n(t) = \sum_k \exp(ikn)s_k(t)$ can be used to yield an equation for the mode amplitudes s_k , *i.e.*,

$$\ddot{s}_k + \gamma \dot{s}_k + \omega_k^2 s_k - 4au_0 \cos(\omega t + \theta_0) s_k = 0, \quad (8)$$

where we discarded a higher harmonics and introduced $\omega_k^2 = \omega_0^2 + 4\kappa \sin^2(k/2) - 8(au_0/\omega_0)^2$. Setting $\tau = \omega t/2$ and $s_k(t) = v_k(t) \exp(-\gamma t/2)$, one derives a Mathieu equation

$$\ddot{v}_k + [A - 2Q \cos(2t + 2\theta_0)] v_k = 0, \quad (9)$$

with the parameters $A = (2\omega_k/\omega)^2 - (\gamma/\omega)^2$ and $Q = 8au_0/\omega^2$. If it holds that $\sqrt{A} \simeq l$, with l denoting a positive integer number, the Mathieu equation allows for parametric resonance [32,33]. The extension of the resonance regions is determined by the ratio Q/A ; for the primary resonance, $A \simeq 1$, it is given by

$$(A-1)^2 < Q^2. \quad (10)$$

For the parameter set corresponding to the line shown in the inset in fig. 1 (determining the relationship between the amplitude f of the external ac-field and the amplitude of the homogeneous solution u_0) the instability bands for different values of the coupling strength κ are depicted in fig. 2. For the onset of parametric resonance the driving amplitude f has to exceed the value of the bifurcation point, *i.e.* $f_c \gtrsim 0.1408$ related with the “jump” resonance, regardless of the value of κ . The position of the bottom of the instability band, determining the critical unstable wave number k_c , shifts towards lower k values with increasing coupling strength κ . For a chosen field strength $f=0.15$, that lies just above f_c , one expects that the LMs

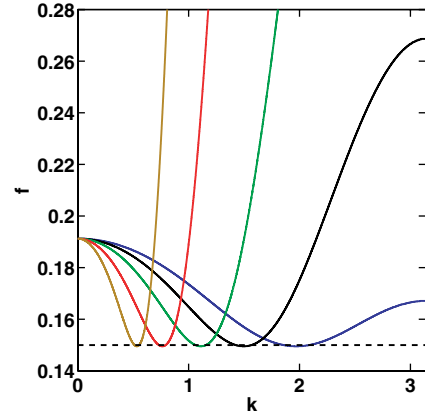


Fig. 2: (Color online) Instability bands for different coupling strengths $\kappa = 2, 1, 0.5, 0.3, 0.2$ (decreasing from left to right) are displayed. The relation between the amplitude of the external ac-field f and the amplitude of the homogeneous solution u_0 is the one displayed with the inset in fig. 1. The horizontal dashed line at $f = 0.15$ intersects each instability band very close to its bottom, the position of which determines the respective critical wave number k_c (see also text).

of distinct wavelength, determined by $\lambda_c = 2\pi/k_c$ become excited (cf. fig. 2). We infer from fig. 2 that the wavelength of a LM increases with increasing coupling κ . This is verified in fig. 3 showing the spatio-temporal evolution of the amplitudes $q_n(t)$ for couplings $\kappa = 0.5$ and $\kappa = 2$. The Langevin equations were numerically integrated using a two-step Heun stochastic solver. In all our simulations the initial chain configuration is represented by $q_n(0) = x(0)$ and $p_n(0) = 0$ with the homogeneous solution (plane wave) $x(0)$ given in (7). We note the formation of a LM of certain wavelength arising from the homogeneous state after a short time span (after $t \sim 60$) and we note that the period duration for oscillations near the bottom of the potential is around $2\pi/\omega_0 \simeq 4.4$.

Regarding the energy relation within the stationary flat state (where each unit contains the same amount of initial energy) we monitored the temporal evolution of the energy of one unit

$$E_n = \frac{1}{2} p_n^2 + U(q_n), \quad (11)$$

and the corresponding field energy

$$E_{field} = -f \sin(\omega t + \theta_0) q_n, \quad (12)$$

without coupling the chain to the heat bath for a force amplitude $f = 0.15$. In this stationary case the field energy performs small-amplitude oscillations around a mean value of $E_{field} = 0.04 \equiv 0.03 \times \Delta E = 0.03 \times 4/3$, while the mean of the energy of one unit is $E_n = 0.78 = 0.585 \times \Delta E$ (not shown). Thus the gain of energy, determined by the ratio E_n/E_{field} , amounts to a remarkable high value of 19.5. To retain this relation upon lowering (increasing) the damping γ a lower (higher) driving strength f is necessary while the “jump” resonance frequency attains a lower (higher) value according to eq. (6).

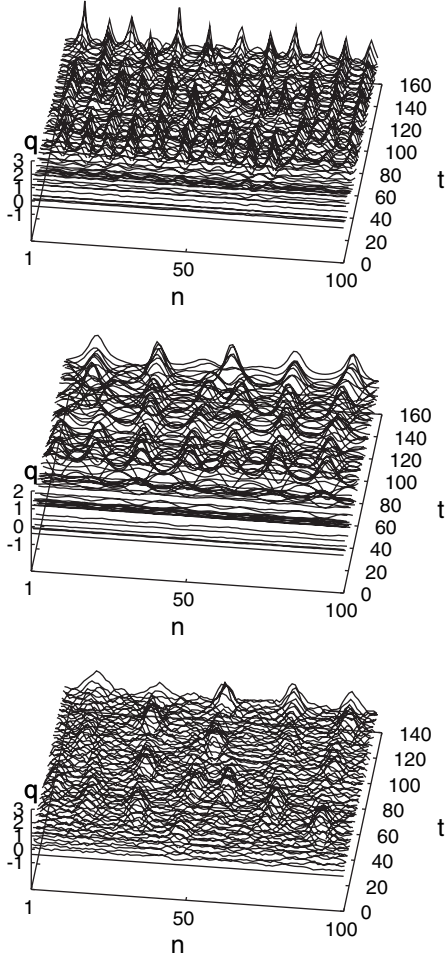


Fig. 3: Spatio-temporal pattern of the solutions $q_n(t)$ for a lattice consisting of $N = 100$ sites and chosen coupling strength at $\kappa = 0.5$ (top panel), $\kappa = 2$ (central panel) for the same realization of Gaussian white noise with thermal energy $k_B T = 0.001 \times \Delta E$. Bottom panel: Same as in the central panel but now for a 50-times-larger thermal energy: $k_B T = 0.05 \times \Delta E$. The remaining parameter values are set at $f = 0.15$, $\omega = 1.295$, $\theta_0 = 0$ and friction $\gamma = 0.1$.

The stochastic term provides perturbations of all wave numbers and a pattern emerges from the homogeneous flat state. That is, due to the effect of parametric resonance perturbations provided by the thermal noise grow and induce a LM consisting of several humps. The fastest growing perturbations are those associated with the critical wave number k_c (see also [34]). Each of these humps resembles the hairpin shape of the transition state as the critical escape configuration possessing an energy E_{act} through which the coupled units have to pass in order to cross the barrier [30]. The robustness of the LMs is remarkable: a LM is sustained, despite continuously impacting thermal noise of strengths up to values $k_B T \lesssim 0.2 \times \Delta E$. Moreover, the formed pattern maintain their distinct wavelength $\lambda_c = 2\pi/k_c$ (see fig. 3).

We note that upon increasing the noise strength the growth rate of the humps becomes enhanced, being

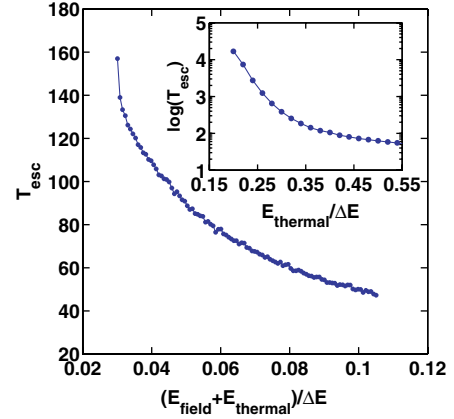


Fig. 4: The mean escape time of the chain *vs.* the mean injected energy $E = E_{field} + k_B T$ measured in units of ΔE with *fixed* field energy $E_{field} = 0.03 \times \Delta E$ provided by an external modulation field with $\omega = 1.295$, $\theta_0 = 0$ and $f = 0.15$. Here we vary the thermal energy $E_{thermal} = k_B T$. The inset depicts the unforced case with $f = 0$. The remaining parameter values are $N = 100$, $\kappa = 0.28$ and $\gamma = 0.1$.

reflected in the statistics of the barrier crossing of the chain in the presence of weak ac-driving. The amplitude and frequency of the latter are chosen such that the dynamics exhibits parametric resonance. The dependence of the mean escape time of the chain on the injected average energy $E \equiv E_{field} + E_{thermal}$, with $E_{thermal} \equiv k_B T$ (measured in units of the barrier energy ΔE) is displayed in fig. 4. The thermal energy $E_{thermal}$, supplied non-coherently by the heat bath, is *varied* within the range $[(10^{-4} - 0.11) \times \Delta E]$.

The average of the escape times was performed over 500 realizations of the thermal noise. In this context the random escape time of a unit is defined as the time instant when the unit passes through the value $q = 20$ far beyond the potential barrier. Thus, no likely recrossing back into the potential valley can occur [29,30]. The escape time of the chain is then determined by the average of the escape times of its units. We notice that the underlying irregular dynamics serves for self-averaging and thus the choice of the phase of the coherent, external forcing, θ_0 , does not affect the mean escape time. In the forced as well as unforced case there occurs a rather rapid decay of T_{esc} with growing $E_{thermal} = k_B T$ at low temperatures. This effect weakens gradually upon further increasing $k_B T$. Most strikingly, for the forced system the escape times become drastically shortened in comparison with the unforced case with $f = 0$. Moreover, for the forced system escape takes place also at very low temperatures for which in the undriven case not even the escape of a single unit has been observed during the simulation time (taken here as $t = 10^5$) implying a *giant enhancement of the rate of escape* as compared to the purely thermal-noise-driven rate.

Upon exploring the optimal escape route, we investigated the influence of the coupling strength κ on the

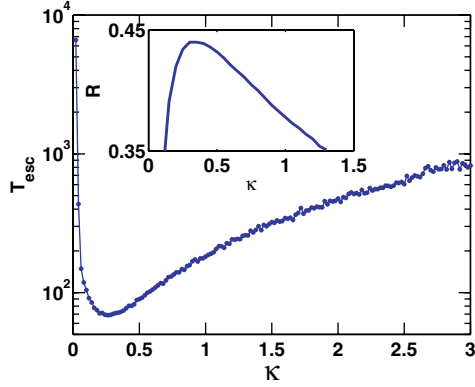


Fig. 5: The mean escape time *vs.* the coupling strength κ exhibits a resonance structure. The inset displays the ratio R , defined in eq. (13), as a function of κ . The remaining parameter values are given by $N = 100$, $k_B T = 0.05 \times \Delta E$, $f = 0.15$, $\omega = 1.295$, $\theta_0 = 0$ and $\gamma = 0.1$.

average escape time. Our numerical findings are summarized in fig. 5. The mean escape time exhibits a *resonance structure*, viz. there exists an optimal coupling strength ($\kappa_{res} \simeq 0.28$) for which the escape assumes a minimum. Upon lowering $\kappa < \kappa_{res}$ we notice a drastic rise of the escape time while for $\kappa > \kappa_{res}$ the graph exhibits only a moderately growing slope with growing coupling strength κ . We emphasize the *collective* nature of this resonance effect which here occurs for *finite* interaction strength $\kappa \neq 0$. In the limit $\kappa \rightarrow 0$ the mean escape time of noninteracting, individual particles assumes for this parameter set an extreme large value, implying a vanishingly small escape rate.

To explain the occurrence of the resonance structure in fig. 5 we recall that the wavelength, $\lambda_c = 2\pi/k_c$, of the arising LMs on the lattice is determined by the critical wave number $k_c = k_c(\kappa, f)$ (cf. fig. 2). The number of humps contained in a LM, N_h , can be attributed to k_c as: $\lambda_c N_h = 2\pi/k_c N_h = N$. The number of humps (besides their height and width) regulates how the mean energy injected via the coherent external field and the incoherent thermal noise is shared among them. Supposing that the whole lattice can be divided into an array of segments, where each of them supports a single localized hump, the energy of one segment is given by $E_s = E/N_h = 2\pi E/(k_c N)$. Appropriate conditions for successful escape are provided when the energy contained in each segment, E_s , is close to the activation energy, E_{act} of the critical escape configuration [30]. The efficiency of energy localization is then determined by the ratio

$$R = E_s/E_{act}. \quad (13)$$

The activation energy as a function of the coupling strength satisfies (we recall that we use a dimensionless formulation) the relation $E_{act} = (1 + 3.54 \times \kappa)\Delta E$ [30]. Keeping the injected energy E fixed and given value of k_c we obtain R . In the inset of fig. 5 the ratio R is plotted

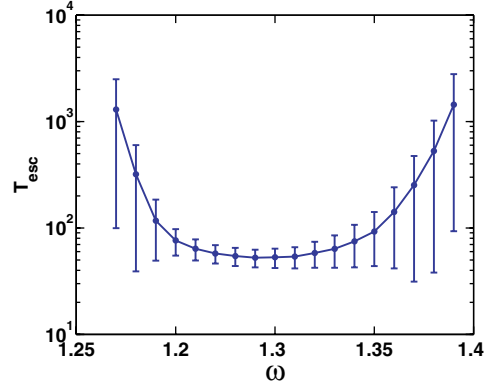


Fig. 6: The mean escape time as a function of the driving frequency. The thermal energy is $k_B T = 0.05 \times \Delta E$ and the driving amplitude is $f = 0.15$. The remaining parameter values are $N = 100$, $\omega = 1.295$, $\kappa = 0.3$, $f = 0.15$ and $\gamma = 0.1$. For comparison we note that Langer’s theory [3] yields for the mean escape time in the unforced case, $f = 0$, the value $T_{esc} \simeq 553 \times 10^5$ underpinning the drastic speed-up effect in our forced case.

as a function of the coupling strength κ . The plot indeed exhibits a maximum at $\kappa = 0.28$, which confirms the finding of the resonance found for the mean escape time *vs.* coupling strength as depicted in fig. 5. Concerning the critical localized mode through which a lattice state has to pass through in order to escape over the potential barrier, we remark that for comparatively low coupling strengths ($\kappa \lesssim 0.6$) the effect of discreteness in the lattice system is still so pronounced that this critical localized mode is indeed represented by a thin hairpin-shaped configuration, involving effectively one lattice unit of large amplitude to either side of which the amplitude pattern decays extremely rapidly (for more details see [30]).

Next we study the role of the angular driving frequency of the external modulation field, see in fig. 6. The escape time as a function of the angular frequency likewise exhibits a resonance structure and there exists an optimal frequency for which the average escape time assumes a minimum. This is reminiscent of the phenomenon of resonant activation found for the thermally activated escape of noninteracting particles surmounting oscillating barriers [23–27]. In our case of a nonlinear chain composed of coupled units, however, this “resonant activation” within a frequency window nicely correlates with the systems’ gain of energy that is coherently supplied by the applied ac-field in this very same frequency interval (note the corresponding frequency window in the nonlinear frequency response graph associated with large amplitudes in fig. 1). Therefore, tuning the frequency ω at a fixed interaction strength κ allows to optimize further the mean escape time. The minimal escape scenario thus requires an optimal tuning both in coupling strength and ac-driving frequency ω .

In summary, we have presented a drastic speed-up mechanism of the thermal-noise-driven barrier crossing

of a coupled damped nonlinear oscillator chain under the impact of a weak external ac-field. With appropriate parameter values of the latter and in the presence of thermal noise an instability mechanism is initiated due to which LMs arise from stationary flat-state solutions in the lattice dynamics. Humps of the LMs are rapidly driven through the transition state thus accelerating the escape over the situation with purely thermally assisted escape. Interestingly, the LMs are sustained up to fairly high noise levels corresponding to $k_B T \simeq 0.2 \times \Delta E$.

The findings of our study can be applied for the control of the rate of barrier crossing of oscillator chains. With such chains providing the archetype model for nonlinear collective transport of matter, charge and energy in abundant low-dimensional systems in physics, biology and chemistry, this speed-up scenario of thermally driven collective escape over potential barriers might well be put to constructive use in a variety of potential applications.

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