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Sound and Noisy Light: Optical Control of Phonons in Photo-switchable Structures

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We present a novel means of controlling phonons via optical tuning. Taking as a model an array of photo-responsive materials (photoswitches) embedded in a matrix, we numerically analyze the vibrational response of an array of bistable harmonic oscillators with stochastic spring constants. Changing the intensity of light incident on the lattice directly controls the composition of the lattice and therefore the speed of sound. Furthermore, modulation of the phonon bandstructure at high frequencies results in a strong confinement of phonons. The applications of this regime for phonon wave-guides, vibrational energy storage, and phononic transistors is examined.

Light-matter interactions are of great utility for many of applications. For mechanical vibrations (phonons) these applications encompass creating or destroying phonons using light (optomechanics [1, 2], Raman scattering [3–7]), creating or destroying light using phonons (vibronic spectroscopy [8], black body radiation [4]), and shifting the refractive index using phonons (acousto-optics [9–11]). However, the last quadrant of this interaction (Table I), the control of phonon properties (e.g. speed of sound, \(c_s\)) using light remains unexplored. As a result, acousto-optic devices (filters, modulators) are common, whereas controlling phonons remains difficult. Fundamentally, phononic devices require more than the manipulation of phonon populations, but also the tunable manipulation of speed and transmission (phase and amplitude) [12–14]. Optical control, a fast, non-contact technique, is a natural candidate for this. The absence of existing optical control methods for phonons is surprising, as it is well-known both that light can tune other material properties (particularly magnetic and electronic properties) by means of inducing a structural phase transitions (a.k.a. nonlinear phononics, ionic Raman) [5–7, 15–19], and that other signals (pressure, temperature) can control \(c_s\) and phonon dispersion (often by phase transitions) [12–14, 20–29]. So far as we are aware, the only research that came close to the problem of optical control were [30–32]. [30] showed that the vibrational properties of a material can be optically switched in photo-responsive liquid crystal polymers. But the optical excitation was intense enough for complete switching, so their focus was rather on the thermal and strain (rather than the optical) tuning of this switching. Whereas [31] considered fs laser pulses on bismuth, where photo-induced thermal expansion induces an optical fluence dependent red-shift in some of the phonon modes. The effect is limited by the system melting, as it’s an expected signature of materials near a phase transition [33]. It is also similar to the earlier work of [32] on chalcogenide glasses, where the photo-softening was associated with the approach of a glass-liquid melting transition. There have been other scenarios where optical driving has affected phonon dynamics, such as [34], but these have been switches between discrete phases (and therefore not tunable) or changed particular resonant modes (i.e. polaritons).

<table>
<thead>
<tr>
<th>Photon (\rightarrow) Phonon</th>
<th>Create/Destroy</th>
<th>Harden/Soften</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optomechanics, Raman</td>
<td>???</td>
<td></td>
</tr>
<tr>
<td>Vibronic, Fluorescence</td>
<td>Acousto-Optics</td>
<td></td>
</tr>
</tbody>
</table>

TABLE I: The quadrants of light-matter interaction. First row is light used to control phonons, second is sound used to control phonons. First column is changing populations, second is changing dispersion.

generic photo-switches and demonstrate its use at modulating the phonon bandstructure, tuning \(c_s\). Numerical analysis of the dynamics reveals additional confinement effects at high-frequencies which may prove useful for vibrational energy storage or phononic transistors.

Consider a solid under illumination, there exist three photon-phonon coupling mechanisms. If the basis atoms are polar, then there is a coupling to the electromagnetic field inducing a localized vibration (direct Raman). If there exists an infrared-active phonon mode, then photons can excite that (ionic Raman). Absent these couplings, light can excite electrons, which excite phonons via electron-phonon coupling (indirect or stimulated Raman). After this initial excitation of a single mode, nonlinearities (electron-phonon coupling or phonon anharmonicity) will disperse this energy into a thermal phonon population. For some materials, these excitations can also induce a structural change, driving the vibrations about some new equilibrium position in ionic Raman or softening a phonon mode in stimulated Raman. Structural changes necessarily change the phonon bandstructure. To avoid conflating the creation of thermalized phonons and the tuning of the band-structure (distinguishable effects for sufficiently weak anharmonicity, [35]), we concentrate on non-photoactive modes and neglect thermalization except for its effect in structural transitions. For consistency, this requires neglecting phonon-phonon couplings so we concentrate on harmonic phonons. Hence we confine attention to a chain of 1D simple harmonic oscillators (SHOs) without further loss of generality.

We therefore model a system of \(n\) masses \((m \equiv 1)\) joined to \(N = n + 1\) photo-switchable SHOs (ground state spring constant \(k_D\), excited \(k_E\)) with a mechanical driving force \(F = F_0 \cos(\omega_0 t)\) at frequency \(\omega_0\). Driving pumps a constant supply of energy into the system, so it is helpful to
modify the standard clamped boundary conditions \( u_0 = 0 = u_1 \), \( u \) is displacement) by sandwiching this system between impedance matched systems of \( n \) damped (damping rate \( \gamma_k \equiv 1 \)) SHOs \( (k = k_D) \) and clamping these ends (see Fig. 1). In the more general case, exciting any SHO will not produce switching elsewhere (i.e. no cascades). This is plausible for sufficiently separated photo-isomerizing molecules and composite or multilayered structures where only some portion is photo-sensitive (see the bottom of Fig. 1). However, if there are no photoswitching cascades, then the order and timing of (de)excitation greatly influence the dynamics. To avoid a biased order, switching is randomized with Poisson statistics (excitation rate \( R_D = B_{D U} H \) and de-excitation \( R_U = A_{spont} + B_{U D} H \) where \( H \) is photon fluence at a point and \( A \) and \( B \) are the Einstein coefficients [4]). This is plausible in the case of sufficiently low intensity photo-excitation that shot noise dominates (i.e. individual photon trajectories are relevant, not an ensemble of photons) but is also a technique for ensuring the robustness of the response to changes in the switching order. Switching dynamics are typically complicated, but because the photo-excitation is much faster than any structural rearrangement, these complexities can be neglected (to lowest order) by integrating out the shorter time-scales to give

\[
\dot{k}(t) = -\gamma_k(k(t) - k_{SS}(w))
\]

where \( \gamma_k \) determines the rate of the structural reaction (typically \( O(\omega_i) \)) and \( k_{SS} \) is the new steady-state, \( w \) denotes the stochastic variable describing switching. In principle a change in the equilibrium position of the lattice is also possible. But our system models a photo-switch embedded in a matrix and does not describe the (realization-dependent) dynamics of modes shorter than one supercell, so this shift is negligible. Hence, the displacement obeys

\[
\ddot{u}_i(t) = k_i(t) [u_{i-1}(t) - u_i(t)] + k_{i+1}(t) [u_{i+1}(t) - u_i(t)]
\]

where \( i \) indexes the site.

At steady-state this system’s eigenmode distribution is solvable using Random Matrix Theory [36]. However, this does not describe the effects of changing composition (i.e. traversing the RMT’s solution space [37]). Moreover, the eigenfunctions for a single equation can be solved – exponential wave-functions when no switching is occurring and modified Bessel functions of imaginary order when switching is occurring – but the inconsistency of this basis set impedes an analytic solution for any non-trivial realization. As such, we integrate the solutions numerically. Switching times/locations, being stochastic, are computed using the Gillespie algorithm [38], and the system can be integrated analytically between switchings. Initial conditions are \( u(x, 0) = 0 = \dot{u}(x, 0) \), \( k_i(0) = k_D \). We use natural units: \( a \) the equilibrium site separation for length and \( t = 1/\gamma_0 \) for time. Using \( k_D = 1/t^2, k_U = 2/t^2, R_U = 1.5/t_{\gamma}, R_D = 2/t_{\gamma}, F_0 = 1a/t^2 \), a sample of size \( n = 29 \) is calculated for interval \( T = 100t_{\gamma} \) giving \( u(x, t) \) and \( k(x, t) \) at various \( \omega_0 \).

![FIG. 1: Schematic model of system. Photoswitches (green) sandwiched between damped regions (brown) which are clamped at the far ends. Light (orange curves) is applied to the photoswitches, which are also driven mechanically (blue arrow) to produce phonons. Photoswitches are modeled as a series of 1D SHOs composed of photosensitive materials (e.g. anthracene, see bottom) embedded in a matrix, creating a bistable system with two spring constants.](image-url)
ing the dispersion and thereby $c_s$. To generate the dispersion we Fourier transform $u(x, t)$, giving $u(q, \omega)$. The location of the maxima of $u(q, \omega)$ indicates the mode $\omega(q)$ that was excited by driving at $\omega_0$. (To generate a smooth dispersion rather than a series of discrete normal modes, $n = 124$ is used.) This is repeated for $N_{\text{rep}} = 10$ times with $T = 200t_\gamma$ for ergodicity, and $\omega$ and $q$ are averaged. Because at high frequencies waves can be narrowly confined, this will artificially introduce Fourier components near the $\Gamma$ point. Since these are not features of the waveform itself, but of its confinement, we exclude these terms from the average. Repeating this for multiple frequencies allows us to construct the dispersion, which is repeated for various combinations of $R_U/R_D$ in Fig. 3.

Note that for points above $\omega_q$ (i.e. in the confinement regime), some of the dispersions show a pronounced drop in $q$. This is an artifact of the exponential decay regions, which lower the effective wavelength. Ergo, this jump is a result of limitations in defining a wavelength for such a heterogeneous system. So when we fit the dispersion $\omega(q; R_U/R_D)$ to a sine curve $c_{s, \text{eff}} \sin(a_{\text{eff}} q)/a_{\text{eff}}$ (the dispersion for a homogeneous 1D chain, $c_{s, \text{eff}}$ and $a_{\text{eff}}$ are fitting parameters) it is helpful to exclude these points.

From the fitted dispersion we can extract the effective $c_{s, \text{eff}}$ by expanding for small $q$ giving (in units of $a/t_\gamma$)

$$c_{s, \text{eff}} = 0.997 + 0.177 N_U/N + 0.227 (N_U/N)^2$$

(3)

where $N_U$ is the excited state population, $c_{s, D} = 1a/t_\gamma$ is the ground state’s speed of sound and $c_{s, U} = \sqrt{2a/t_\gamma}$ is the excited state’s (see Fig. 3 inset). From the master equation

$$\dot{N}_D = -R_D N_D + R_U N_U = -\dot{N}_U$$

(4)

the steady state compositions are

$$\frac{N_U}{N} = \frac{R_D}{R_D + R_U} = \frac{(g_U/g_D) H}{S(\omega_\nu) + (1 + g_U/g_D) H}$$

(5)

where $g_U/D$ are the mode degeneracies, $S = 2\hbar \omega_\nu^3/\pi e^3$, and $\omega_\nu$ is the photon frequency (last relation comes from detailed

FIG. 2: Phonon amplitude under photo-switching. $x$-axis denotes distance along the 1D chain, $y$-axis denotes time. Contours (coloured) are isoval curves of phonon amplitude $u_i(t)$ (see sidebar). Grey segments are the ground state (no illumination), white segments are excited (illuminated), gradients are transitions. (a) Low frequency ($\omega = 0.5/t_\gamma$). Bottom inset (red, vertical slice) shows $u(N/2, t)$ (red, solid curve) and $k(N/2, t), k(N/2 + 1, t)$ (black, dashed curves) as a function of time. Side inset (blue, horizontal slice) shows a snapshot $u(x, t_0)$ (blue, solid curve) and $k(x, t_0)$ (black dots) as a function of position. (b) Mid frequency ($\omega = 1.5/t_\gamma$). (c) High frequency ($\omega = 2.5/t_\gamma$).

FIG. 3: dispersion $\omega(q)$ for a 1D chain of photoswitches. Color denotes strength of illumination from blue (no illumination, all in ground state) through red (full illumination, all in excited state). Dotted lines and dots (with error bars) indicate numerical results. Solid lines denote fit to sinusoid dispersion. (inset) Speed of sound as a function of fraction in the excited state. Colored, solid line is the function of $q$, $\nu$ denotes distance along the 1D chain, $u$ [arb. u.]

$\omega = 0.5, R_{\uparrow} = 1.5, R_{\downarrow} = 2, k_D = 1, k_U = 2$. Time ID:735650.2972

$\nu = 1.5, R_{\uparrow} = 1, k = 1/t_\gamma^2$.
balance of the Einstein coefficients [4]). Changing $R_U / R_D$ changes $N_U / N_D$, so increasing the optical intensity increases the equilibrium population in the excited state (to some limiting fraction given by the mode degeneracies). Ergo, changing illumination gives direct control of $c_s$. Now consider a simple kinetic model where $c_s$ in the ground (excited) state is $c_{s,D} (c_{s,U})$. The average speed for an inhomogeneous system would be the weighted average $c_{s,D} N_D / N + c_{s,U} N_U / N$ or
\[
e^{(kin)} c_s = c_{s,D} + (c_{s,U} - c_{s,D}) N_U / N \quad (6)
\]
i.e. linear with composition. This disagrees with our observed relation, which falls below this kinetic limit except for the homogeneous cases of $N_U / N = 0$ or 1 which agree with the analytic results to within 99\% accuracy (see Fig. 3 inset). This is expected, though, as reflections delay a pulse and decrease its effective velocity.

Finally, consider again the confined regime. In the homogeneous cases the transmittivity of the material should be nearly 1 (i.e. no loss) or 0 (i.e. perfect damping) for a sufficiently thick sample. Switching between these compositions (possible when $g_U > g_D$ and photon intensity is large, or stochastically possibly for $R_D > R_U$ and $N \approx g_U / g_D$) allows for illumination controlled switching between transmission and reflection. Dynamically changing $R_D / R_U$ therefore allows for controlled phonon transmission. This switching mechanism is therefore a potential phononic transistor using the optical analog indirect control scheme presented in [14] (i.e. a light source instead replaces the electromagnet). Such an indirect transistor is more easily tuned than the direct designs, which rely upon phonon-phonon couplings [29] that are not dynamically accessible. To show the feasibility of this proposal, we repeat our calculations for a pulsed illumination ($R_D^{(on)} = 4 / \gamma$, $R_D^{(off)} = 0$). Pulse widths are selected such that a homogeneous composition is produced for each state. Since complete, monotonic switching of a sample has the expectation value
\[
R_U, D^{[N]} = N \sum_{1}^{N} 1 / \gamma = N H_N \approx N \ln N \quad (7)
\]
where $H_N$ is the harmonic function, we use a sample size of $n = 9$ for these simulations (illumination period $200 \tau_x$, dark period $200 \tau_x$, total run time 2000 $\tau_x$, $R_D = 4 / \gamma = 8 R_U$). Plotting the amplitude at the far end of the sample (normalized to the maximum amplitude without damping $\max[u_0(L,t)]$) gives Fig. 4 for $\omega = 2.1 / \tau_x$. The horizontal lines indicate the rms average amplitude during each period of darkness or illumination (including switching intervals, therein underestimating the difference). Frequencies below $\omega_2$ there is transmission for both states so the ratio of these averages is nearly 1 (not shown). Whereas for frequencies just above $\omega_2$ there is a large difference between the states and so a large separation observed (Fig. 4). As frequency increases above $\omega_2$, transmission drops and the ratio again approaches 1 (not shown). Comparing these results with Fig. 2c reveals a cross-over from confinement to transmission with increasing photon intensity. For confinement to be effective, there should be narrow domains of propagating configuration, which is best achieved with weak driving.

In summary, we have demonstrated a novel approach controlling the phononic properties of a system, using light to tune the phonon band structure. This reverses the acousto-optic formulation of phonons modulating the index of refraction, and instead light modulates $c_s$. The shifting of the dispersion that this allows opens several interesting new possibilities for phononic devices. Delay lines and by extension phase control gates can be constructed by tuning the speed of sound. Thermal conductivity modulation is clearly achievable by the controlled scattering of short-wavelength phonons from the configuration boundaries. Vibrational energy storage or phononic memory are possible in the high frequency regime with phonon confinement under weak optical driving. And under the same regime with strong optical driving, an opto-phononic switch or indirect phononic transistor is feasible. The control of the speed of sound could also improve the short-term storage (RAM) of phononic information through delay line memory, similar to [39]. Furthermore, the patterning of photoswitches in a system — or their patterned photoexcitation — allows for real-time, adaptable phononic materials, including phononic crystals and metamaterials. This can be particularly useful (in the confinement regime) for creating waveguides that dynamically control the propagation of phonons or for tuning the thermal conductivity by selectively introducing scatterers. Such an approach would also be experimentally feasible, as an optically-controlled bandpass filter was proposed using a superlattice of resonant cavities with chalcogenide glasses in [40] (only light or dark states were simulated and so no intensity-dependant tunability was observed). Similar approaches for experimentally realizing these devices could be achieved by superlattices of photo-sensitive materials (e.g. photo-isomers [30], ionic Raman active materials [31], photo-elastic glasses like chalcogenides [32]) separated by inactive layers or by embedding these photo-sensitive
materials (particularly photo-isomers, which are generally small organic molecules) within an inactive matrix (the scenario illustrated in Fig. 1). Finally, The prevalence of phonon couplings in quantum computing, electronics, photonics, and spintronics [41–45] implies that these effects may have further applications in the optical control of a great many signals in a cavalcade of fields. Thus, our “opto-phononic” framework introduces an unexplored aspect of light-matter interaction beyond the previously studied thermal repopulation regime.

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[35] Thermalization and velocity modulation are distinct effects, assuming sufficiently weak nonlinearity to the lattice. Considering a current of phonons \( J = n v \), thermalization will change the number of phonons \( n \), while structural changes will change the velocity \( v \). It is only when anharmonicity is so strong that the band structure is intensity dependent that this distinction breaks down. And even then, the use of low amplitude phonons makes this negligible.
[37] Changing compositions is also equivalent to traversing different realizations of a glassy or disordered medium, or taking an ensemble average over disorder.