Phonon Bloch oscillations in acoustic-cavity structures

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We describe a semiconductor multilayer structure based in acoustic phonon cavities and achievable with MBE technology, designed to display acoustic phonon Bloch oscillations. We show that forward and backscattering Raman spectra give a direct measure of the created phononic Wannier-Stark ladder. We also discuss the use of femtosecond laser impulsions for the generation and direct probe of the induced phonon Bloch oscillations. We propose a gedanken experiment based in an integrated phonon source-structure-detector device, and we present calculations of pump and probe time dependent optical reflectivity that evidence temporal beatings in agreement with the Wannier-Stark ladder energy splitting.

Electronic Bloch oscillations, that is, oscillations of an electron induced by a constant electric field in the presence of a periodic potential, are a beautiful and clear example of quantum effects in solids. When an electric field is applied on a charged particle in a crystal, its wavevector increases with time. Thereafter, Bragg interference leads to a velocity reduction, and finally to a sign change at the band edge. Notwithstanding its simplicity, for many years the issue was controversial and only quite recently the existence of electronic Bloch oscillations has been definitively established. In normal metals the large Brillouin zone and electron relaxation lead to an overdamped behavior characterized by Ohm’s law. Instead, electronic Bloch oscillations are observable in SL’s due to the Brillouin zone reduction.

Very recently photon Bloch oscillation devices based in optical microcavities have been proposed, and related reflectivity and time-resolved optical transmission features have been observed in structures grown on porous silicon. The concept behind these devices is also quite nice and simple. The optical microcavities provide the discrete photonic states which can couple through photon mirrors thus leading to photonic minibands. The spatial dependent energy gradient, equivalent to the electric field for electrons, can be achieved, e.g., by varying the refractive index. As compared to electrons, photon dephasing mechanisms are less effective and thus Bloch oscillations are, in principle, more easy to observe. On the other hand, photonic minibands require a large number of optical microcavities (around 30-50), amounting to total thicknesses larger than 40-50 microns. This is too large for current molecular beam epitaxy (MBE) semiconductor technology. For this reason the reported structures were grown using electrochemical methods of porous silicon nanostructuring that do not have this limitation. The drawback is that the quality of the samples is not as good, layer interfaces and refractive indices are not that well controlled, and optical residual absorption becomes an important issue in the performance of the structures.

Sound in solid media is also described by a wave equation, the relevant parameters being the material density and sound velocity, and the boundary conditions establishing the connection between displacement and strain between two different materials. Extending the above concepts applied to electrons and photons to phononic structures, in this letter we propose semiconductor multilayer structures capable of displaying acoustic phonon Bloch oscillations. The building block of the structures are acoustic phonon cavities as recently introduced by Trigo and coworkers. The phonon wavelength is only a few nanometers (phonon frequencies being in the THz range), and thus the sample full size and required interface layer quality are achievable with actual MBE technology. In addition, acoustic phonon mean free paths are large compared to the structure size (typically below a micron) and thus dephasing is not a critical issue. We present calculations of phonon reflectivity, displacement distribution, and time evolution upon excitation with a localized phonon source to illustrate the device behavior and to demonstrate the existence of a phonon Wannier-Stark ladder (WSL) and Bloch oscillations (BO). In addition, using a photoelastic model we calculate the Raman spectra which we show give a direct measure of the created phononic WSL. Phonon BO can be independently probed using coherent phonon generation techniques. For this purpose we propose a specifically designed device and we present calculations of time dependent reflectivity induced by femtosecond impulsions in this structure.

A periodic stack of two materials with contrasting acoustic impedances reflects sound. The first k = 0 folded phonon mini-gap in a SL is maximum with the layers thickness (d) ratio given by d₁/v₁ = d₂/3v₂, the stop-band and reflectivity of such a phonon mirror being determined by the acoustic impedance mismatch \( Z = \rho₁ v₁ / \rho₂ v₂ \) and the number of SL periods. A phonon cavity can be constructed by enclosing between two SL’s a spacer of thickness \( d_c = m\lambda_c / 2 \), where \( \lambda_c \) is the acoustical phonon wavelength at the center of the phonon minigap. The cavity confined modes correspond to discrete energy states within the phonon stop-
a broad stop-band (between $\sim 15$ and $\sim 21.5$ cm$^{-1}$), basically determined by the superposition of the individual cavity minigaps. A series of dips modulate the reflectivity within this wide stop-band. Between the lower stop-band limit and $\sim 17.4$ cm$^{-1}$, and between $\sim 18.6$ cm$^{-1}$ and the upper stop-band limit, a series of features with varying energy spacing can be identified. These originate from phonon interferences determined by a propagation limited by the sample surface and by the minigap bands. These features are relatively weak and broad because of the small acoustic impedance mismatch between the top GaAs layer and the cavity structure. On the other hand, sharp reflection dips are observed in the region between $\sim 17.4$ and $\sim 18.6$ cm$^{-1}$. These dips, which are equidistant and separated by $\Delta = 0.15$ cm$^{-1}$ correspond to the coupling of external phonons, by tunneling through the minigap band, to the Wannier-Stark states confined within the structure. (iii) This phononic WSL can be also identified by the well defined discrete phonon modes displayed in panel (c). The WSL is the spectral domain counterpart of the BO. It is precisely in this spectral region where oscillations should appear in the time-domain.

The time and spatial variation of the acoustic phonon displacement field $U_g(z,t)$, created by a pulse described by a spectral function $g(\omega)$ and incident at $t = 0$ at the GaAs-sample interface ($z = 0$), can be evaluated using the scattering method described by Malpuech and Kavokin. Within this description, $U_g(z,t) = \frac{1}{\pi} \int_{-\infty}^{\infty} u(z)g(\omega)\exp(-i\omega t)\,d\omega$, where $u(z)$ are the stationary solutions of the elastic wave equation with frequency $\omega$ shown in Fig. 1(c). The time evolution of such wavepackets for the two different energies indicated in Fig. 1(a) are shown in Fig. 2. For energies above or below the phonon WSL region (20 cm$^{-1}$ in the example shown), the incident pulse propagates within the sample up to a position where it is back-reflected by a minigap band leaving afterwards the sample. On the contrary, when $\omega$ corresponds to the WSL energy region (18.3 cm$^{-1}$ in the displayed figure), a fraction of the pulse energy is backreflected at the surface while another part enters the structure by tunneling through the minigap band developing afterwards clear periodic oscillations within the structure. In order for these phonon BO to be observed, the FWHM of $g(\omega)$ should be larger than $\Delta$. For these calculations we have used a gaussian distribution with $2\sigma = 1.0$ cm$^{-1}$. On the other hand, the period of the oscillations (and consequently the length travelled by the pulse) is directly determined by $\Delta$. ($\tau_B = h/\Delta$).

In what follows we will show how Raman scattering and coherent phonon generation experiments can provide a direct probe of the phonon WSL and BO, respectively, in these devices. Raman scattering has extensively shown to be a powerful tool to study phonons in semiconductor multilayers and, in particular, confined modes in acoustic cavities. In order to evaluate the Raman spectra, we use a photoelastic model. We analyze two experimental geometries, namely backscattering ($BS, k_S \sim -k_L, q = \sim 2k_L$) and forward scattering ($FS, k_S \sim k_L, q \sim 0$). Here $k$ refers either to the
scattered or laser wavevector, and \( q \) is the transferred wavenumber. In Fig.3 we present BS and FS spectra calculated for the sample described above and assuming laser excitation with 550nm. For comparison purposes the corresponding phonon reflectivity is also shown. The Raman spectra display a complex series of peaks in the stop-band spectral region. We have verified that such rich spectra are a kind of sample finger-print that can be used as a characterization tool. Interestingly, the BS and FS spectra display clear peaks and dips, respectively, at exactly the WSL energies. High resolution Raman setups working in the visible can discern spectral features with resolution better than 0.02 cm\(^{-1}\), thus providing a spectral-domain tool able to probe the underlying phonon WSL.

Coherent phonon generation is termed the impulsive generation of phonons using high power ultrashort laser pulses.\(^{11}\) In the case of THz vibrations, femtosecond pulses are required. To the best of our knowledge the generation mechanism is still an open issue, and no complete theory is available to describe the processes involved. We briefly describe next a model for pump and probe coherent phonon generation and detection based on a photoelastic coupling between light and acoustic phonons.\(^{12}\)

This mechanism is the only active when pump and probe are below the gap. Above the gap other mechanisms can contribute, but we expect the main conclusions to remain essentially valid. Any arbitrary time and position dependent phonon displacement in the structure \( w(z, t) \) can be expressed in terms of the phonon eigenstates as

\[
w(z, t) = \int r_\omega u(\omega, z) sin(\omega t) d\omega.
\]

Assuming that phonons are generated coherently through a photoelastic mechanism by a femtosecond pulse (modelled as \( E_0(z, t) \propto \delta(t) exp(ik_z z) \)), the coefficients for the above expansion can be obtained as

\[
r_\omega = \frac{2}{L} \int_0^L p(z) |E_0(z)|^2 \frac{\partial u(z)}{\partial z} dz.
\]

Here \( L \) is the length of the sample, \( p(z) \) is the photoelastic constant which is assumed constant in each layer, and \( \partial u(z)/\partial z \) is the strain associated with an eigenstate of energy \( \omega \). We note that the FS (\( g = 0 \)) Raman cross section discussed above is proportional to \( |r_\omega|^2 \). Once this excitation is generated, it evolves according to the time dependence \( sin(\omega t) \) and can be detected by a delayed, lower power, probe pulse that senses the time dependent change of reflectivity.\(^{13}\) The change in reflectivity can be calculated as

\[
\Delta R(t) \propto \int_0^\infty \Delta \omega(z, t) exp(2ik_z z) dz,
\]

where the probe pulse has been assumed to be proportional to \( e^{ik_z z} \). Introducing again the photoelastic coefficient to relate \( \Delta \omega \) with the strain \( \delta w(z, t)/\partial z \), the time dependent change in reflectivity can be written as

\[
\Delta R(t) \propto \int_0^\infty p(z) \frac{\partial w(z, t)}{\partial z} exp(2ik_z z) dz.
\]

If \( p(z) \) is constant, this equation implies that only phonons with \( q = 2k \) can be probed. On the other hand, the detector’s \( p(z) \) can be designed to access other excitations by backfolding the phonon dispersion. Comparing this equation with the Raman cross section,\(^{14}\) it is easy to see that the observable phonon spectrum is related to the BS Raman scattering component of the detector structure.

In order to generate the quasi-monoenergetic phonons required to induce BO, and to monitor the time evolution of the latter, we have conceived a monolithic source-sample-detector device (see the scheme in Fig. 4). The first GaAs/AlAs SL (SL/s in Fig. 4) acts as the phonon source and is designed to generate, by excitation with a fs impulsion above the gap, an elastic pulse with energy centered at 18.3 cm\(^{-1}\) and width equal to \(~ 0.8 \text{ cm}^{-1}\).\(^{11}\) It is made of 30 periods of 43.2 Å/51.5 Å GaAs/AlAs. The layer widths determine the energy of the generated pulse, while the number of periods define, due to finite size effects, its width. The coefficient \( r_\omega \) of the phonon pulse (or equivalently the FS Raman spectra) generated in this SL is shown in the inset on Fig. 4. Once generated, the coherent phonons propagate into the structure and act as the phonon source \( g(\omega) \) for exciting Bloch oscillations. Between the SL source and the structure, a GaAs 200nm buffer layer is introduced to screen the residual pump power, not absorbed in the SL, from impinging into the cavity structure and generating unwanted frequencies. The \( g(\omega) \) phonon pulse, on the other hand, propagates through the GaAs layer basically unaltered. The cavity structure is identical to the one described above. Once within this structure, Bloch oscillations develop and part of the energy is lost to the substrate upon each turn. Their effect at the right end side of the sample is probed by the second SL (SL/d in Fig. 4), which acts as an energy selective detector of the Bloch oscillations through their effect in the time variation of the reflectivity.\(^{12}\) To keep this time variation simple, a second GaAs layer is introduced to stop the probe beam from being modulated also by the cavity structure. The BS Raman spectrum of the SL defines the detector’s bandwidth, which we have designed to include the pulse \( g(\omega) \). It consists of a 20 periods 38.1 Å/68.1 Å GaAs/AlAs SL. Its BS spectrum, calculated for a 750 nm probe pulse, is shown in the inset on Fig. 4. The device is terminated by a thick (> 1µm) Ga0.5Al0.5As layer which acts both as a stop layer for chemically etching the GaAs substrate, and as a window for accessing the detector SL from the back. In Fig. 4 we present calculations of the reflectivity change as a function of time. Fast oscillations dominate the reflectivity, corresponding to the frequency of the coherently excited phonons (18.3 cm\(^{-1}\)). In addition, this fast component is amplitude modulated by an envelope whose frequency is determined precisely by the Bloch oscillation period. The Fourier transform of the time dependent reflectivity clearly shows the WSL frequencies within an envelope determined by the input source \( g(\omega) \) and the detector bandwidth.

In conclusion we have extended concepts previously discussed in the context of electronic and photonic properties of solids to acoustic phonon physics. We have described semiconductor structures based in recently
reported acoustic cavities and achievable with actual growth technologies, displaying phononic Stark ladders and capable of sustaining Bloch oscillations. We have also shown how Raman scattering and coherent phonon generation provide spectral and time domain probes, respectively, of these acoustic phenomena. Structures as the one described here can be exploited to enhance the coupling between sound and other excitations (electrons and photons), and as the feedback high-Q resonator of a phonon laser. Moreover, engineered phonon potentials are not limited to linear dependencies that mimic electric fields but can take arbitrary shapes. This opens the way to novel phonon devices based in the discrete confined states of acoustic cavities.

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FIG. 1: (a) Phonon band structure. The central white region limited by the thicker black forbidden bands corresponds to the cavity-mode miniband. The white and thinner black regions above and below this bands arise from Bragg oscillations. The arrows indicate the energy of the pulses displayed in Fig. 2. (b) Phonon reflectivity. In (a) and (b) shaded areas indicate the WSL region. (c) Phonon displacement as a function of position and energy. Darker regions indicate larger acoustic phonon intensities. The arrows label the WSL energies. The solid lines limit the “forbidden” bands. See text for details.

FIG. 2: $\log(u(z,t))$ for gaussian phonon wavepackets centered at the two energies indicated with arrows in Fig. 1(a), and FWHM given by $2\sigma = 1 \text{ cm}^{-1}$.

FIG. 3: Back and forward 550 nm Raman scattering spectra corresponding to the acoustic cavity structure with $N_c = 25$ described in the text. For comparison the corresponding phonon reflectivity is also shown. The inset highlights the Stark ladder spectral region (17-19.5 cm$^{-1}$).

FIG. 4: Top-left: Source and detector SL’s Raman spectra. Top-right: Scheme of the proposed monolithic source-sample-detector phonon device (see text for details). Bottom-left: Time variation of the reflectivity probed at the detector SL. Bottom-right: Fourier transform of the detected reflectivity change.
