Inclusion of Bandstructure and Many-Body Effects in a Quantum Well Laser Simulator

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A self-consistent eight band $k\cdot p$ calculation, which takes into account strain and includes Hartree, exchange, and correlation terms (determined from a local density approximation) is incorporated into a QW laser simulator (MINILASE-II). The computation is performed within the envelope function approximation for a superlattice, in which all spatially varying terms of the $k\cdot p$ Hamiltonian, including the exchange and correlation energies are expanded in plane waves. The $k\cdot p$ eigenvalue equation, and Poisson’s equation are solved iteratively until self-consistency is attained. Results from the $k\cdot p$ calculation are exported to MINILASE-II via a density of states and an energy dependent optical matrix element factor, renormalized by a Coulomb enhancement factor to account for electron-hole attraction. Results are presented for the gain spectrum and modulation response for a Ga$_{0.8}$In$_{0.2}$As/Al$_{0.1}$Ga$_{0.9}$As quantum well laser with and without the inclusion of the Coulomb enhancement factor.

Keywords: Gain, $k\cdot p$, laser, quantum well, modulation response, bandstructure

MINILASE-II [1] is a semiconductor laser simulator that self-consistently solves Poisson’s equation, the carrier transport equations (both drift diffusion and thermionic emission across heterostructures), quantum well (QW) capture, and the photon rate equations for arbitrary two dimensional geometries. Many important effects arise from the bandstructure near the quantum well. However, it is computationally impractical to perform an accurate self-consistent bandstructure calculation within the laser simulator at each iteration. The purpose of this paper is to explain how we connect a separate eight band $k\cdot p$ superlattice calculation, including many-body effects, with MINILASE-II and to present results for the gain and modulation response for a strained-layer In$_{0.2}$Ga$_{0.8}$As/Al$_{0.1}$Ga$_{0.9}$As quantum well laser.

The $k\cdot p$ calculation involves diagonalizing a Hamiltonian which at the center of the Brillouin zone can be expressed as

$$H_{nl}^{(kp)}(k_{||}, k_z) = \left( E_n(0) + \frac{\hbar^2 (k_{||}^2 + k_z^2)}{2m} \right) \delta_{nl}$$

$$+ (k_{||} + k_z \hat{e_z}) \cdot \mathbf{P}_{nl}$$

$$+ \sum_{\alpha \beta} \Gamma_{\alpha \beta}^{nl} k_{\alpha} k_{\beta} + H^{(strain)}$$

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where the indices run over the bands in our basis set $B$, $P_{\nu\ell}$ are the momentum matrix elements, and $\Gamma_{\nu\ell}^{(e)}$ are renormalization constants describing the contribution from bands not contained in $B$. To characterize the bandstructure of crystals with zincblende symmetry near $\Gamma$, it is sufficient to include the heavy hole (HH) and light hole bands (LH) ($\Gamma_{11}^0$), the split-off bands (SO) ($\Gamma_{5}^0$) and the lowest conduction bands ($\Gamma_{6}^0$). $H^{(\text{strain})}$, a $k$-independent term, describes the strain, which is assumed to be confined to the well region. The resulting $8 \times 8$ Hamiltonian has been described in the literature [2, 3].

For the superlattice calculation, $z$ is taken to be the growth direction and the usual substitution $k_z \rightarrow \partial_z / i$ is made to obtain the effective mass Eq. [4],

$$\sum_{\ell \in B} H_{m\ell}(k||, \partial_z / i) \phi_{m}(n_{k||, k_\ell})(z) = E^{(n)}(n_{k||, k_\ell})(z), \quad (2)$$

where $k_\ell$ is the wave number for the 1D Brillouin zone of the superlattice, and $\phi_{m}(n_{k||, k_\ell})(z)$, the superlattice envelope functions, are related to the wave functions to second order by

$$\psi_{m||} = \exp(i k_{||} \cdot r) \sum_{n \in B} \left( u_{m0} + \sum_{\ell \in B} \frac{k_{||} \cdot P_{\ell m}}{E_{\ell}(0) - E_{m}(0)} u_{\ell 0} \right) \phi_{m}(n_{k||, k_\ell})(z). \quad (3)$$

The band parameters which enter the Hamiltonian are allowed to have different values in the well and the barrier regions [5], and the resulting operator is Fourier transformed. For a superlattice period of 500 Å, typically about thirty Fourier components are required to ensure convergence of the ground state eigenenergies to within 1 µeV. The figures throughout this paper were generated for a superlattice consisting of Al$_{0.1}$Ga$_{0.9}$As barriers of 400 Å, and In$_{0.2}$Ga$_{0.8}$As wells of 80 Å. The barriers are wide enough to decouple adjacent wells so that the superlattice is in practice a collection of independent quantum wells. Most band parameters were determined from [6] except for the conduction band and valence band deformation potentials used to determine band edge shifts due to strain and the fraction conduction band discontinuity, which are not well known. We assume a fraction conduction band discontinuity of 0.7 and 11 eV for the difference in conduction band and valence band deformation potentials.

To account for carrier–carrier interactions additional terms which depend on electron and hole densities are added as diagonal terms in the $k \cdot p$ Hamiltonian. The direct Coulomb interaction gives the Hartree potential $V_H$ which is determined by solving the 1D Poisson equation:

$$\frac{\partial_z}{\partial_z} (\varepsilon \partial_z V_H) = -\frac{4\pi}{\varepsilon} \rho_{\text{3D}}(z) \quad (4)$$

$$\rho_{\text{3D}}(z) = \frac{\rho_{\text{2D}}(L)}{L} \int_{BZ} \frac{d^2k}{(2\pi)^2} \sum_{j} \left| \phi_j(k, z) \right|^2 f(E_j(k), \mu(n_{2D})) \quad (5)$$

where the envelope functions $\phi_j$ have been normalized to the superlattice period $L$. The Fermi distribution $f(E, \mu)$ depends on the energy dispersions found from the $k \cdot p$ calculation and on the quasi-chemical potential $\mu$, determined from the input parameter $n_{2D}$, the carrier density per unit surface in one period of the superlattice.

$$n_{2D} = \sum_{j} \int_{BZ} \frac{d^2k}{(2\pi)^2} f(E_j(k), \mu) \quad (6)$$

The interaction of a carrier with its exchange-correlation hole lowers its energy and results in a narrowing of the bandgap. This effect is taken into account in the local density approximation. We use an interpolated expression derived by Hedin and Lundquist [7] for three dimensional systems, and treat electrons and holes as separate plasmas. The bandgap renormalization (BGR) obtained is in qualitative agreement with experiment and can be improved by using an expression more appropriate for 2D systems.

Because these additional terms depend on the eigenfunctions determined from the $k \cdot p$ calcula-
tion, an iterative approach must be used. First the \( \mathbf{k} \cdot \mathbf{p} \) Hamiltonian is diagonalized for a suitable initial guess for the Hartree, exchange and correlation energies. For a given 2D carrier density, the chemical potentials are then computed assuming a Fermi distribution function. The \( z \) dependence of the 3D carrier density can then be determined and used to compute a new estimate of the Hartree and exchange and correlation potentials. To assure convergence, an underrelaxation scheme is used for the exchange and correlation potentials. The approach is illustrated in the flow chart in Figure 1.

Once convergence has been attained, optical matrix elements (OME) given by \( M_{\nu\nu'} = \langle \psi_{\mathbf{r}k}\mathbf{p} \mathbf{r} | \psi_{\mathbf{r}'k}\mathbf{p} \mathbf{r}' \rangle \) are computed for later use in MINILASE II. The square of the OME gives the strength of the electron-hole coupling through the photon interaction and therefore enters the expressions for spontaneous and stimulated emission. Figure 2a shows the OME dispersion for TE polarization and two directions of \( \mathbf{k}_\parallel \), parallel to and perpendicular to \( \hat{\mathbf{e}} \). At \( \Gamma \), CB1 couples only to HH1 and LH1 because of the orthogonality relation between the envelope functions. However, away from \( \Gamma \), band mixing allows formerly forbidden transitions to take place. As expected for this polarization, the coupling at \( \Gamma \) between CB1 and HH1 is about three times stronger than between CB1 and LH1 [8].

Due to the Coulomb attraction between electrons and holes, spontaneous and stimulated emission are enhanced. This Coulomb enhancement, which may be thought of a renormalization of the optical matrix elements, is computed external to MINILASE II for various carrier densities and lasing energies. The expression used for the Coulomb enhancement factor has been

\[
H_{\text{Coul}} = \frac{\mu^2}{\epsilon_{\text{vac}}} \left( \frac{2a}{2\pi} \right)^3
\]

FIGURE 1  Flowchart for self-consistent \( \mathbf{k} \cdot \mathbf{p} \) calculation.

FIGURE 2  (a) Optical matrix elements for \( k_{\parallel} \) parallel to (solid) and normal to (dashed) the polarization. Transitions between the lowest conduction subband CB1 and the valence subbands HH1, HH2 and LH1 are shown. (b) Density dependence of the Coulomb enhancement factor.
derived in the literature [9] and is given by

\[
\frac{1}{1 - q(k)}
\]

where

\[
q(k) = \frac{1}{M(k)} \int dk' V^{(s)}(k - k')
\]

\[
\frac{iM(k')}{\hbar} \frac{f_n(k') + f_p(k') - 1}{i(E_n(k') - E_v(k') - \nu) + 1/\tau}
\]

Here \( f_n \) and \( f_p \) are Fermi occupation factors, \( \tau \) is the carrier-carrier scattering time, assumed to be 100 fs, and \( V^{(s)} \) is the 2D screened Coulomb potential which in the static long wavelength limit of RPA is given by

\[
V^{(s)}(q) = \frac{2\pi e^2}{\varepsilon_0 q} \frac{1}{1 + \kappa/q}
\]

where

\[
\kappa = \frac{2\pi e^2}{\varepsilon_0} \left( \frac{\partial n}{\partial \mu_n} + \frac{\partial p}{\partial \mu_p} \right)
\]

is the inverse screening length. Figure 2b shows the renormalization factor for the CB1-HH1 transition between the lowest conduction subband and lowest valence subband for several carrier densities above threshold. In this calculation, an axial average of the matrix elements is performed, so that the OME may be treated strictly as a function of energy rather than in-plane wave vector \( k_{||} \). We see that the modest enhancement of the OME decreases as the carrier density increases. This result stems from the fact that screening is enhanced at higher densities.

The results of these calculations are exported to MINILASE II in three forms: the spatial profile of the envelope functions, a density of states and an optical matrix element factor \( P_n(\Omega) \) (associated with each conduction band energy grid point \( n \)) given by

\[
P_n(\Omega) = \sum_q \int \frac{d^2 k_{||}}{(2\pi)^2} |\hat{\mathbf{e}} \cdot \mathbf{M}(k_{||})|^2 \eta_n(E_n'(k_{||}))
\]

\[
\delta(E_n'(k_{||}) - E_v'(k_{||}) - \Omega)
\]

The direction of the light polarization is \( \hat{\mathbf{e}} \), and \( \eta_n \) is a hat function with support on the energy range defined by grid points \( n-1 \) and \( n+1 \) and whose area is normalized to unity. The sum extends over all conduction subbands \( i \) and valence subbands \( j \).

The gain coefficient is then given by

\[
g^{(0)}(\Omega) \sim \frac{1}{\hbar \Omega} \sum_n \left( \tilde{f}_e(\varepsilon_n) + \tilde{f}_h(\varepsilon_n - \Omega) - 1 \right)
\]

\[
P_n(\Omega) \mathcal{L}(\Omega, \tau)
\]

where \( \mathcal{L} \) is a Lorentzian broadening function and \( \tilde{f}_e \) and \( \tilde{f}_h \) are the non-equilibrium electron and hole distribution functions, respectively. Minilase II incorporates the k-p data in an iterative manner. That is, the simulator first solves the carrier transport equations, Poisson's equation and the photon rate equations through Newton's method. The resulting carrier densities are used to recompute the band edges from an interpolated expression for the bandgap renormalization. Then, files appropriate for the computed density are read from disk. Since the exchange-correlation terms are added as k-independent diagonal elements, their inclusion does not significantly alter the effective masses and therefore the densities of state, except for shifting the band-edges.

We now briefly describe a few results obtained from MINILASE-II for an operating regime beyond the lasing threshold. Figure 3(a) shows the optical gain spectra with and without the Coulomb enhancement factor for the same applied current. In steady state, the maximum height of the gain is pinned by the losses in the laser which is the same in each case. There are two salient features that distinguish the gain curve without the Coulomb enhancement from the other curve. These are the reduced threshold for the onset of gain and the larger transparency point. Both features stem from the smaller effective matrix element in the absence of Coulomb enhancement.
that requires greater pumping of the laser to achieve a fixed gain. Thus, more carriers are required which results in a reduction in the band gap and implies a greater separation of the electron and hole quasi-fermilevels and thus a larger transparency point.

Figure 3(b) displays the laser's response to a small square pulse in the applied voltage. We note that the frequency peak is blue shifted when a larger effective matrix element is used. This may be explained as follows. The photon rate equation has the form

\[ \dot{S}_\nu = (G_\nu - 1/\tau)S_\nu + R \]

where \( \nu \) is a mode index, \( S_\nu \) is the photon population, \( G_\nu \) is the modal gain, \( \tau_\gamma \) is the photon lifetime and \( R \) is the spontaneous emission rate, which is much smaller than the other terms of the right hand side under lasing conditions. Since the losses \( 1/\tau \) are fixed, the gain is roughly proportional to the time derivative of the photon density in the cavity which implies a more swiftly responding laser.

In conclusion, we described how an accurate \( k \cdot p \) calculation was connected to a sophisticated laser simulator and demonstrated how the larger effective matrix element that stems from the Coulomb enhancement affects the gain and resonance frequency in the modulation response of a laser.

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