Photocarrier extraction in GaAsSb/GaAsN type-II QW superlattice solar cells

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Photocarrier transport and extraction in GaAsSb/GaAsN type-II quantum well superlattices are investigated by means of inelastic quantum transport calculations based on the non-equilibrium Green’s function formalism. Evaluation of the local density of states and the spectral current flow enables the identification of different regimes for carrier localization, transport, and extraction as a function of configurational parameters. These include the number of periods, the thicknesses of the individual layers in one period, the built-in electric field, and the temperature of operation. The results for the carrier extraction efficiency are related to experimental data for different symmetric GaAsSb/GaAsN type-II quantum well superlattice solar cell devices and provide a qualitative explanation for the experimentally observed dependence of photovoltaic device performance on the period thickness. Published by AIP Publishing. https://doi.org/10.1063/1.5030625

Multi-junction solar cells enable photovoltaic solar energy conversion efficiencies beyond 46%1 due to enhanced utilization of the solar spectrum and the reduction of thermalization losses.2,3 Maximization of the efficiency requires suitable combination of materials with specific band gap energies. Among the most promising candidates are 4-junction devices based on (Al)InGaP and (In)GaAs as top cells, where the challenge is to find a third cell with the band gap energy of 1–1.15 eV that is lattice matched to (In)GaAs and the Ge substrate.4 Recently, strain-balanced GaAsSb/GaAsN quantum well superlattice (QWSL) structures have been proposed as high-quality subcell materials with tunable bandgap close to the target range.5,6 In addition to the absence of clustering7 due to spatial separation of N and Sb, the type-II SL (SL-II) offers largely independent control of conduction and valence band edges via the individual adjustment of the N and Sb content, respectively.8 The tunability of the effective bandgap by variation of the SL period thickness d_per was confirmed by photoreflectance measurements for d_per ∈ {3, 6, 12, 20} nm, showing effective band gap energies in good agreement with theoretical predictions from SL dispersion calculations based on effective masses and band offsets derived from the double band-anticrossing (2-BAC) model.9,10 In order to move towards low effective band gaps, larger period thicknesses are favored in principle. However, as displayed in Fig. 1, the external quantum efficiency (EQE) experiments revealed serious performance issues related to incomplete carrier extraction at short circuit conditions for d_per = 12 nm (SL-II12) as compared to the situation for d_per = 6 nm (SL-II6), where the coincidence of the EQE at short circuit and at large reverse bias points at unit extraction efficiency.9 Hence, the SL period thickness has a critical impact on the photovoltaic device performance, which requires careful analysis in order to identify optimum configurations.

Here, we perform such an analysis by means of rigorous simulation of photocarrier transport in GaAsSb/GaAsN QWSL structures with varying thicknesses of the GaAsSb and GaAsN layers. Our approach relies on steady-state quantum-kinetic theory in the shape of the non-equilibrium Green’s function formalism (NEGF) as adopted for the description of photovoltaic device operation in nanostructure-based solar cell devices.11,12 As in previous applications of NEGF to the simulation of QWSL solar cells,13,14 the electronic structure is modeled in terms of a simple effective mass Hamiltonian describing decoupled conduction and valence bands. Planar geometry is assumed, with the in-plane spatial coordinates Fourier-transformed to transverse momentum \( k_{\parallel} \). In the present case, the effective masses of the bulk ternaries and the band offsets defining the SL heterostructure potential along the growth direction \( z \) are obtained from the 2-BAC model. While this amounts to a crude simplification of the complex electronic structure of dilute nitrides and does not allow for quantitatively accurate computations in an extended spectral range, it provides a reasonable qualitative explanation for the experimentally observed dependence of photovoltaic device performance on the period thickness.

FIG. 1. Measured external quantum efficiency for GaAsSb/GaAsN QWSL solar cells with period thicknesses of 12 nm (SL-II12) and 6 nm (SL-II6), respectively. Comparison of the data at short circuit (\( V = 0 \) V) and at large reverse bias (\( V = -3 \) V), where all the photogenerated carriers are extracted, reveals a massive deterioration of the extraction efficiency for the thick period, which points at configuration-dependent transport issues.

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description of the states occupied via optical transitions at photon energies close to the effective bandgap, as considered here. Table I displays the parameters for the N content $x_N = 1.2\%$ and the Sb concentration $y_{Sb} = 3.25\%$, as experimentally determined for the samples of Ref. 5. This configuration with low N and Sb contents is nearly lattice-matched to GaAs and exhibits very low levels of strain in the constituent layers; the effects of strain are therefore neglected in the model. The corresponding band offsets given by the 2-BAC model are $\Delta E_C = 194\text{ meV}$ and $\Delta E_V = 50\text{ meV}$ [cf. Fig. 2(a)]. Any other material parameters required for the quantification of the interaction of carriers with photons and phonons, i.e., the parametrization of the respective interaction Hamiltonians, are chosen as of bulk GaAs.

Conventionally, the properties of SL structures are analyzed based on models assuming perfect periodicity and flat band conditions. The appropriateness of both assumptions for the finite systems under consideration will be assessed in the following. Figure 2(b) displays the one-dimensional density of electron states (at vanishing transverse momentum, $k \parallel = 0$) of the GaAsSb/GaAsN type-II SL with $d_{\text{per}} = 6\text{ nm}$ obtained by integrating the local density of states (LDOS) from the Green’s functions over the entire finite structure, $\mathcal{D}_{1D}(E) = -L_z^{-1} \int dz \text{G}^R(0, z, z, E)/\pi$. The 1D density of states (1D-DOS) is shown for the increasing number of SL periods, revealing a fast convergence of the miniband width, but features related to the finite size of the structure remain up to very large systems. For comparison, we also plot the 1D-DOS of the perfectly periodic SL, which is evaluated by $\mathcal{D}_{1D}(E) = 2 \left\{ \partial_{q_0} \text{e}_n(q_0) \right\}^{-1} / \pi |q_0| (z = E)$ using the longitudinal SL dispersion $\text{e}_n(q_0)$ for $n = 1$ as obtained from solving the Schrödinger equation for the same effective mass Hamiltonian but with periodic boundary conditions. There is a close agreement between the finite and infinite system DOS for the band width; however, some discrepancies remain in shape and asymmetry of the DOS inside the band.

In order to investigate localization and transport of photogenerated charge carriers in finite SL structures with different period thicknesses, we evaluate the (1D) LDOS—corresponding to the integrand of the formula for the 1D-DOS—and the spectral current density $j(E) = \mp (2\varepsilon \hbar/\pi) \lim_{\zeta \to z} \left( \delta_{z, \zeta} - \delta_{\zeta, z} \right) \int dk |G^S(k, z, \zeta)|^2 / (2\pi)^3$, where the upper (lower) sign is for electrons (holes), for the quasi-ballistic case, i.e., in the presence of elastic scattering only. In Fig. 3(a), the result is displayed for a SL-II 6 structure with 12 periods and carrier-selective contacts as

| GaAsSb$_{0.0325}$ | GaAsN$_{0.012}$ |
|------------------|------------------|
| el. affinity $\chi$ (eV) | 4.076 | 4.270 |
| Band gap $E_g$ (eV) | 1.364 | 1.220 |
| el. effective mass $m^*_e/m_0$ | 0.067 | 0.098 |
| hl. effective mass $m^*_h/m_0$ | 0.516 | 0.500 |

FIG. 3. Local density of states (for $k_\parallel = 0$) and (normalized) spectral current density flow under monochromatic illumination for (a) SL-II 6 (at $E_c = 1.3\text{ eV}$) and (b) SL-II 12 (at $E_c = 1.25\text{ eV}$) configurations in the absence of built-in fields and inelastic scattering. In both cases, the states are delocalized over the entire SL and extraction proceeds quasi-ballistically at the energy of generation, with no indication of a thickness-dependent change in the qualitative behavior.

FIG. 2. (a) Schematic layer structure and the band profile of the GaAsSb/GaAsN type-II SL considered in this work. (b) Evolution of the 1D density of states (at $k_\parallel = 0$) in the first miniband of the SL-II 6 configuration with the increasing number of periods. For comparison, the SL potential and the DOS of the infinite (perfectly periodic) SL are shown as well. While the miniband width is converged already at 15 periods, finite-size effects remain as small features in the DOS up to very large period numbers.

### Table I. Bulk parameters for constituent materials used in the simulations as derived from the double band-anticrossing model$^{9,10}$

| Material | $E_g$ (eV) | $\chi$ (eV) | $\mu$ (eV) | $\mu_0$ (eV) |
|----------|-----------|-------------|------------|--------------|
| GaAs     | 4.076     | 3.469       | 0.866      | 0.752        |
| Sb       | 4.270     | 3.469       | 0.866      | 0.752        |
required for photovoltaic device operation. The LDOS shows clear miniband formation and wave function delocalization over the entire structure for the electrons. In the case of the holes, coupling of periods is weaker but still visible. The quasi-ballistic current under monochromatic illumination over the entire structure for the electrons. In the case shows clear miniband formation and wave function delocalization features. At \( d_{\text{per}} = 3 \text{ nm} \), both electron and hole states are delocalized over many periods; at \( d_{\text{per}} = 6 \text{ nm} \), holes have become localized, while electrons are still delocalized; and at \( d_{\text{per}} = 12 \text{ nm} \), the lowest states of both carrier species are localized.

Figure 4 shows the electron and hole 1D-LDOS at a built-in field of 15 kV/cm—obtained from a drift-diffusion-Poisson simulation of the full experimental \( p-i-n \) structures described in Ref. 5—and at short-circuit conditions (\( V_{\text{bias}} = 0 \text{ V} \)). For \( d_{\text{per}} = 3 \text{ nm} \), both electrons and holes are delocalized over multiple periods. At \( d_{\text{per}} = 6 \text{ nm} \), the hole states are confined to a single period, while the electron wave functions are still delocalized. Finally, at \( d_{\text{per}} = 12 \text{ nm} \), both electron and hole ground states are fully localized. The thickness-dependent localization behavior is reflected in the transport mechanism for charge carriers, as shown in Fig. 5, displaying the spectral photocarrier flow, again for monochromatic illumination with \( E_c = 1.25 \text{ eV} \). At \( d_{\text{per}} = 3 \text{ nm} \), photocarrier extraction is fast and proceeds quasi-ballistically for both electrons and holes, irrespective of the presence of inelastic electron-phonon scattering. At \( d_{\text{per}} = 6 \text{ nm} \), electron extraction proceeds still quasi-ballistically, while escape of holes by tunneling is no longer fast, which leads to the appearance of a hot carrier current component that is thermally activated by the absorption of optical phonons. At \( d_{\text{per}} = 12 \text{ nm} \), also the electron flow has developed a significant thermal component, while the tunneling of electrons proceeds in a sequential fashion. This transition in the transport mechanism from quasi-ballistic extraction to sequential tunneling with the thermal component has dramatic consequences for the carrier extraction efficiency, as will be shown next.

Formally, the carrier extraction efficiency is defined as the ratio of carrier extraction and generation rates, which can be expressed as \( \eta_{\text{ext}} = J_{\text{sc}}/J_{\text{gen}} \), where \( J_{\text{sc}} \) is the short circuit current and \( J_{\text{gen}} = q \int dz G(z) \) is the generation current obtained from the local generation rate \( G \). For mesoscopic extension of the active device region as considered here, incomplete extraction is expected only for very low carrier lifetime, as is induced by the presence of fast non-radiative recombination channels. As the description of non-radiative recombination as part of a complete NEGF treatment of photovoltaic devices is still at an early stage, and thus only radiative recombination is included here, the low-lifetime regime is made accessible by boosting the rate for spontaneous emission. The lifetime \( \tau \) is then approximated via the
relationship \( R_{\text{rad}} = \delta \rho / \tau \), where \( \delta \rho \) is the photogenerated excess carrier density. In order to study the impact of the period thickness on the photocarrier extraction efficiency, the latter is evaluated for values of \( d_{\text{per}} \) from 3 nm to 12 nm at an effective carrier lifetime of \( \tau \sim 10 \text{ ps} \). To exclude boundary effects due to low bandgap contact layers, a buffer period is inserted on both sides. To obtain results that are relevant for large period number devices (as investigated experimentally), the extraction efficiency is converged with respect to the number of SL periods. The convergence shown in Fig. 6(a) for \( E_c = 1.25 \text{ eV} \) reflects the increasing degree of decoupling of neighboring SL periods with the growing period thickness. Figure 6(b) displaying the converged values as a function of period thickness reveals a dramatic drop in carrier extraction efficiency around \( d_{\text{per}} = 8 \text{ nm} \), which can now be attributed to the transition in the photocarrier transport regime from quasi-ballistic to sequential tunneling accompanied by phonon-mediated escape.

Since thermally activated escape has been identified as an important extraction mechanism at larger period thicknesses, it is instructive to consider the temperature dependence of the extraction efficiency. The results shown in Fig. 7(a) confirm the previously established picture in that close to room temperature, transport at \( d_{\text{per}} = 6 \text{ nm} \) does not rely on thermally activated processes, while such processes play a crucial role at \( d_{\text{per}} = 12 \text{ nm} \). The decrease in extraction efficiency at higher temperatures can be explained by the larger population of phonon modes resulting in increased scattering-induced carrier localization. Figure 7(b) displaying the current spectrum at the contacts for \( T = 250 \text{ K} \) (light color) and \( T = 350 \text{ K} \) (dark color), respectively, shows the transfer of current weight from the tunneling to the thermal component with the increasing temperature. At both temperatures shown, the tunneling component dominates for electrons and the thermal component for holes.

In principle, the thicknesses of GaAsSb and GaAsN layers can be varied independently (under the constraint of observing the strain-balancing condition), which offers a wider parameter space for the design of optimized superlattice solar cell structures. Application of our model and simulation approach to the parameter space spanned by \( d_{\text{GaAsSb/GaAsN}} \in \{2, 3, 4, 5, 6\} \text{ nm} \) identifies configurations with thin GaAsSb and thick GaAsN layers as ideal combination exhibiting high photocarrier extraction efficiency at low effective bandgap values, as can be inferred from Figs. 8(a) and 8(b).

In conclusion, we performed quantum-kinetic simulations of photocarrier generation, transport, and recombination in GaAsSb/GaAsN type-II SL solar cells which shed light on the carrier extraction mechanism in the dependence of the geometrical configuration, built-in field and temperature. The experimental observation of a dramatic drop in carrier extraction efficiency from 6 nm to 12 nm period thickness is explained by the transition in the dominant electron transport regime from quasi-ballistic or band-like transport to sequential tunneling accompanied by phonon-assisted thermal escape. Improved carrier extraction efficiency at similar effective bandgap as in the symmetric 12-nm period structure is predicted for asymmetric configurations with the reduced GaAsSb layer thickness.
The approach demonstrated here may thus serve as a valuable tool in the computational design of superlattice solar cells. In future work, it could for instance be used to explore compositions with effective bandgaps closer to the target value for the application in multijunction devices. For a comprehensive assessment of the impact of SL configuration on the photovoltaic device performance, in addition to the intrinsic absorber under uniform field, the contact regions subject to sizable band bending will need to be considered as well, as they might affect carrier extraction, similar as observed, e.g., in simulations of quantum dot SL absorbers or ultra-thin solar cell architectures.

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FIG. 8. (a) Photocarrier extraction efficiency at $E_c = 1.3$ eV for different thicknesses of the GaAsSb and GaAsN layers in one period. (b) Effective bandgap of the corresponding QWSL structures. In combination, periods with thin GaAsSb and thick GaAsN layers are identified as configurations promising enhanced photovoltaic performance.

The qualitative extraction picture does not depend on the exact value of $s$, as long as $1/s$ is comparable to the rate of carrier escape to the contact reservoirs.