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Control of Electron State Coupling in Asymmetric Ge/GeSi Quantum Wells

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Abstract. Theoretical predictions indicate that the n-type Ge/SiGe multi quantum-well system is the most promising material for the realization of a Si-compatible THz quantum cascade laser operating at room temperature. To advance in this direction, we study both experimentally and theoretically asymmetric coupled multi quantum-well samples based on this material system, that can be considered as the basic building block of a cascade architecture. Extensive structural characterization shows the high material quality of strain-symmetrized structures grown by chemical vapor deposition, down to the ultrathin barrier limit. Moreover, THz absorption spectroscopy measurements supported by theoretical modeling unambiguously demonstrate inter-well coupling and wavefunction tunneling. The agreement between experimental data and simulations allowed us to characterize the tunneling barrier parameters and, in turn, achieve a highly-controlled engineering of the electronic structure in forthcoming unipolar cascade systems based on n-type Ge/SiGe multi quantum-wells.

Keywords: quantum wells, semiconductor heterostructures, quantum tunneling, quantum cascade lasers.

Wavefunction engineering in semiconductor heterostructures has led to the development of quantum electronic and photonic devices based on electron tunneling phenomena, such as quantum cascade lasers (QCL) [1-3] and resonant tunneling diodes (RTD) [4, 5]. At the moment, these widely employed devices are based on compound semiconductors belonging to the III-V elemental group (e.g. GaAs), and most of them exploit electron transport in the conduction band (*n*-type doping), which has a bandstructure considerably simpler than the valence band. **Beyond QCLs, other approaches to terahertz generation employing electron wavefunction tunneling through quantum wells include millimeter wave-pumped superlattices [6] and infrared-pumped quantum fountain structures [7, 8]. Recently,** *n***-type Ge/SiGe heterostructures have been identified as a promising material system to realize CMOS compatible light emitters leveraging on intersubband (ISB) transitions in the THz range and operating at (or close to) room temperature [9-11]. In particular, the recent observation of ISB-transition signatures in the 4–12 THz range, featuring long non-radiative lifetimes of tens of picoseconds at relatively high temperature (~100 K) [8, 12-15], suggests that this material**

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system could enable the fabrication of a SiGe-based THz QCL. Besides the great advantage of compatibility with the standard CMOS technology, the use of Ge/SiGe could also potentially overcome severe limitations of the present THz-QCL technology based on III-V compounds: *i*) the weaker electron-phonon interaction in non-polar semiconductors could increase the device operating temperature, currently limited to ~200 K [16]; *ii*) the absence of a spectral range of forbidden propagation (*Reststrahlen* band) in group-IV semiconductors enables larger emission bandwidth for THz QCLs, well above the current limit of ~5 THz [17]; *iii*) the absence of a strong frequency-dependence of the refractive index in group-IV materials could provide the ideal support for THz frequency combs without dispersion compensation [18-21].

In the pathway towards the realization of a SiGe-based QCL device several issues must be still addressed. The growth has to be optimized in order to achieve i) active epitaxial layers being strain-compensated, relatively thick, and of high crystalline quality; *ii*) a high-degree of reproducibility in the composition profile of the ultrathin barriers, as required to control the electronic spectrum, the tunneling rates, and the electron subband lifetime; iii) sharp heterointerfaces to minimize the detrimental impact of interface roughness (IFR) scattering on the QCL gain [22]. Moreover, the development of reliable theoretical tools to model the ISB electronic dynamics in electrically-biased structures beyond the naive rate equation approach, is another mandatory requirement for an effective design and optimization of the active region stack [23]. This calls for detailed knowledge of the actual material parameters for the SiGe system, which can be only obtained through well-designed experiments, supported by an appropriate framework for the theoretical description, in a multivalley non-polar material system, of the coupling between electronic states in adjacent quantum structures, tunneling rates, and state lifetimes. To this end, a pair of asymmetric coupled quantum wells (ACQWs), being the minimal building block of a QCL architecture, constitutes the ideal exemplar to investigate the dependence of the quantumconfined electron states on the heterostructure properties. By tuning the barrier thickness and the relative width of the two wells, one can modulate the energy of the ISB transitions and the spatial property of the excited electronic states, whose wavefunction can be designed with very different degrees of delocalization over the two wells. Their properties can be investigated by absorption spectroscopy, thanks to the asymmetric character of the coupled wells, which allows ISB transitions from the fundamental level to both the first- and the second-excited state, originating from the interaction of the two levels present in the isolated wells.

After having first observed ISBT in decoupled Ge/SiGe multi-quantum well (MCQW) heterostructures [12-15, 24, 25], in this work we perform the subsequent logical step towards a QCL design, unambiguously demonstrating electronic wavefunction tunneling between adjacent *n*-type strain-compensated Ge/Si_{0.2}Ge_{0.8} ACQWs, grown by chemical vapor deposition (CVD). Using Fourier-transform infrared (FTIR) spectroscopy, coupled to numerical calculations [13], as a probe for the inter-well coupling, we extract relevant information on dipole matrix elements, barrier height, and level broadening due to interface scattering [12, 13, 24-26]. Moreover, thanks to a thorough structural analysis, we demonstrate

that the CVD growth of MQW heterostructures on Si wafers combines high-reproducibility with the feasibility of ultrathin barrier thickness required for tunneling transport, as well as a full control of the in-plane lattice parameter, a mandatory prerequisite to precisely fulfill the strain-symmetrization conditions. It follows that, very thick MQW tunneling structures can be realized avoiding strain relaxation, as required to confine the optical mode in the active region of a QCL operating in the THz range.

Methods

	WL	b	Wt	b∟	D ^{SL} nom	D ^{SL} XRD	X _{Ge}	n 2D
S1	12	2.3	5	21	40.3	41.6	0.87	7.2
S2	13	2.3	5	21	41.3	42.8	0.87	9
S3	11.3	2.3	5	21	39.6	40.6	0.87	7.8
S4	12	3.3	5	21	41.3	42.4	0.81	6
S5	12	4	5	21	42	43	0.81	4.6

Table I: Thickness in nm of the wide (narrow) well $w_L(w_i)$, and of the thin (thick) barrier $b_t(b_L)$ for S1-S5 samples. Nominal (D^{SL}_{nom}) and measured (D^{SL}_{XRD}) superlattice period is also reported; x_{Ge} is the Ge content measured in the thin barrier and the sheet carrier density n_{2D} in units of 10^{11} cm⁻², estimated from FTIR absorption spectra following the procedure reported in Ref. 13.

Five strain-compensated Ge/Si0.2Ge0.8 multi- ACQW samples consisting of a wide and a narrow Ge well separated by a tunneling barrier with thickness w_L, w_t, b_t, respectively, were grown by ultra-high vacuum CVD at 500 °C using ultrapure germane and silane, without carrier gases (see Table I and top of Fig.1). This stack, embedded between thick $Si_{0.2}Ge_{0.8}$ barrier layers (b_L) which avoid coupling between the adjacent modules, has been periodically repeated 20 times. To make ISB transitions in the conduction band observable by FTIR absorption spectroscopy, a 10 nm-wide region at the center of the wide well was co-doped by phosphine resulting in an active phosphorous donor density of $\sim 7.10^{17}$ cm⁻³. A dopant diffusion profile featuring a density reduction of 1 decade/10 nm was measured by secondary ion mass spectroscopy on calibration samples (not shown). The active quantum-well stack was grown on a reverse-graded $Si_{1-x}Ge_x$ virtual substrate (RG-VS) on Ge/Si(001) substrates, ending with a 1.5 µm-thick constant composition Si_{0.13}Ge_{0.87} layer (details on the specific RG-VS can be found in Ref. [12]). The reacting gas pressure was 1.2 mTorr, resulting in a growth rate of 6.5 nm/min for the wells and 4.5 nm/min for the barriers. By using these process parameters, we could carefully control the deposited thickness in the sub-nm range and in a repeatable way, as evidenced by scanning tunneling electron microscopy (STEM) performed in a (Cs)-probe-corrected JEOL JEM ARM200CF. In order to get Z-contrast sensitiveness, STEM images have been acquired with an annular dark-field (ADF) detector. A complementary highresolution x-ray diffraction (XRD) analysis was performed using a Rigaku SmartLab with a rotating anode and line-focus geometry featuring a Ge(400) \times 2 channel cut beam collimator and a Ge(220) \times 2 analyzer crystal.

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Absorption spectra have been measured at a temperature T=10 K in side-illuminated single-pass waveguide configuration with an evacuated FTIR spectrometer (Bruker Vertex 70v), equipped with a He-flow cryostat and an automated lithographic polarizer rotator. We used a surface-plasmon waveguide in which light enters and exits from the side facets of the sample cut at 70° with the growth plane, and the top facet close to the ACQWs has been coated with a metal bilayer (Ti/Au, 10 nm/80 nm). In this configuration, the electric field of the radiation propagating through the QW layers is TM-polarized, and then almost parallel to the ISB dipole. In order to filter-out polarization-independent features not related to ISB transitions, such as those due to the dopant absorption in the Si substrate, we evaluated the dimensionless absorption coefficient $\alpha_{zD}(\omega)$ associated with each QW using the dichroic transmission spectra $T(\omega)=T_{TM}(\omega)/T_{TE}(\omega)$, from which we also estimated the sheet carrier density n_{2D} [13].

To design the ACOWs samples, calculate their energy levels and electron wave-functions, and to simulate the optical spectra we have used a multivalley self-consistent Schrödinger-Poisson solver with parabolic subband dispersion. In this model the band alignments of the L, Δ and Γ conduction minima of a biaxially strained Si_{1-x}Ge_x layer grown on a relaxed Si1-yGey substrate are evaluated according to Ref. 13 where the interest reader can also find the values of the material parameters used in the present calculations. Densities of states for the confined states are calculated in the single band approximation, updating the Ge longitudinal and transverse mass to $m_i=1.59 m_0$ and $m_i=0.093 m_0$, as recently reported in Ref. 27. The electronic states have been calculated selfconsistently in the Schrödinger-Poisson iterative scheme, taking into account ionized impurities and contributions to the Hartree potential from electrons at the L, Δ and Γ valleys. Exchange-correlation effects are included in the local density approximation [28] only for the L electrons, which at low temperature represent the large majority of the carriers. The Fermi level as a function of the lattice temperature is obtained from the spatially resolved 3D donor density. In the evaluation of the optical spectra, the impact of Δ_2 states has been neglected due to their spatial separation from L-states and also because the effect of their presence has never been observed in FTIR data acquired in the past from single QW designed with the excited L subband both in or below the energy interval where Δ_2 states are located [13]. Finally, we notice that, while more advanced computational methods such as density matrix or non-equilibrium Green's functions are certainly mandatory to model electron transport in Ge/SiGe cascade heterostructures, our simple multivalley effective mass approach is sufficient to interpret the ISBT features of the FTIR spectra and to validate the set of material parameters here proposed.

Structural Characterization

In Fig. 1(a), we show a high-resolution ADF-STEM image of a sample, grown for calibration purposes, in which we deposited alternated Si_{0.2}Ge_{0.8} and Ge layers with different thicknesses (dark and bright stripes, respectively). A quantitative analysis of the image clearly indicates the presence of SiGe intermixing at the interfaces, resulting in a

broadening of the well/barrier profile over a length-scale of ~0.8 nm. This effect is obviously more relevant in thin layers (thickness< 2.5 nm), well-matching the results of by Bashir *et al.* on nominally identical Ge/SiGe MQWs [29], and thus leading to a thickness-dependent lowering of the confining potential. Nonetheless, taking this intermixing effect into account, we obtained a reasonable agreement of the measured layers thickness with the nominal one [see inset of Fig. 1(a)]. An upper limit of 0.4 nm for the root mean-square IFR can be derived from the high-resolution analysis of several heterointerfaces.

In Fig. 1(b) we show the ADF-STEM of sample S4, where one can clearly observe the abruptness of the interfaces and thickness reproducibility (within ~2%) of the growth in each of the ACQW modules. The high reproducibility and quality of the heterostructures can be also seen from the XRD data reported in Fig. 1(c), where we show the reciprocal space map (RSM) around the (224) reflections acquired on sample S3. The RSM plot indicates that the entire superstructure is lattice-matched to the GeSi buffer layer [30], which exhibits a stoichiometry of x = 0.87 and a slightly residual tensile strain of $\varepsilon_{\parallel} = 0.20\%$, that can be attributed to the difference of the coefficient of thermal expansion existing between the Si substrate and the RG-VS [31]. The measured in-plane lattice parameter $a_{\parallel} = 5.6373$ Å corresponds to a cubic SiGe layer with an equivalent composition $x_{eq} \sim 0.92$, which, in turns, sets the condition for a fully coherent superlattice. From the superlattice-fringe analysis, we obtained an average Ge composition for the ACQW barriers of sample S4 $x_{Ge} = 0.81 \pm 0.01$ (see Table I). The superlattice periods D^{SL}_{XRD} , calculated for the whole sample set, are also reported in Table I. The values measured by XRD agree with the ADF-STEM results and are equal to the nominal values within 2.5%. The effective Ge content determined from the superlattice period measurement was ~ 0.91, thus confirming the matching condition for strain balancing within the experimental error.



Figure 1: (Top) A schematic diagram of an ACQW module with definition of the relevant thicknesses (see Table I). (a) Z-contrast STEM micrograph of a Ge/Si_{0.2}Ge_{0.8} calibration sample (alloy material appears darker); in the inset, the thickness of the Ge well (red) and SiGe barrier (blue) measured by STEM vs. nominal thickness. The narrowest SiGe barrier is 1.6 nm thick. (b) Z-contrast STEM image of sample S4. (c) XRD-RSM around the (224) reflections of SiGe for sample S3. Scale bars are 10 nm in both panels (a) and (b).

Wavefunction Engineering

The electron wavefunctions in ACQWs are very sensitive to the properties of the tunneling barrier as well as to the energy detuning of the isolated-well levels, which hybridize to form states extending over both the two interacting wells. To disentangle these two effects, in the following we compare ISBT spectra acquired on a sample series featuring the same well-thickness but different b_t (S1,S4,S5) and on a sample series with the same b_t but different w_L (S3,S1,S2). In order to interpret the experimental ISB absorption spectra, we employ a self-consistent multi-valley effective mass model [13], with the measured structural parameters as inputs. As starting point, we calculate in Fig. 2 the fundamental (E_0), first excited (E_1), and second excited (E_2) electron states of asymmetric quantum wells with $w_t = 5$ nm as a function of the wide well width w_L , in the uncoupled ($b_t = 10$ nm) and in the coupled regime ($b_t = 2.3$ nm). In order to take into account the unavoidable Si-Ge intermixing of real samples, x_{Ge} has been set to 0.81 and 0.87 (implying different barrier heights) for the uncoupled and coupled regime, respectively (see Table I). Figure 2(a) displays the energy of the states in the uncoupled regime as a function of w_L : at $w_L \approx 14$ nm, the second level of the wide well aligns to the first level of the narrow well, whose confinement energy remains almost unaffected by the presence of the wide well. For smaller (larger) values of w_L , the first excited level of the un-coupled system E_I has a completely "narrow well" ("wide well") character,

as demonstrated by the square of the electron wavefunctions reported in Figs. 2(b) and 2(c) where electronic levels have been calculated for $w_L = 8$ nm and $w_L = 22$ nm, respectively. The fundamental state E_0 is always confined in the wide well.

Figures 2(d)-2(f) instead display the energy of the states and the square of the electron wavefunctions calculated for $w_L = 8 \text{ nm} [(e)]$ and $w_L = 22 \text{ nm} [(f)]$ in the coupled configuration ($b_t = 2.3 \text{ nm}$). Upon comparing Figs. 2(e) and 2(f) with their counterparts in the uncoupled case, i.e. Figs. 2(b) and 2(c), it is apparent that the wavefunctions of E_1 and E_2 , although featuring the same main character, now have a non-negligible amplitude in both wells, due to the inter-well coupling resulting from quantum tunneling through the barrier. State hybridization between the fundamental level in the narrow well and the first-excited state in the wide well peaks when their energy detuning approaches zero, resulting in the typical level-repulsion behavior evident in Fig. 2(d), from which we estimate, at the crossing point $w_L \approx 14 \text{ nm}$, an anticrossing energy of ~12 meV.



Figure 2: Energy of the fundamental, first-excited, and second-excited electron states (E_0 , E_1 , and E_2 , respectively) calculated for (a) un-coupled, $b_t = 10$ nm and $x_{Ge} = 0.81$, and (d) coupled, $b_t = 2.3$ nm and $x_{Ge} = 0.87$, ACQWs as a function of w_L . Electron energy and squared wavefunction at $w_L=8$ nm and $w_L = 22$ nm for uncoupled (b, c) and coupled (e, f) ACQWs (solid and dotted black curves represent the *L* and Δ_2 band profile, respectively). The potential drop across the ACQW period in plot (b) is related to static charge effects associated to the presence of a dopant tail in the tunneling barrier and in the thin well.

We now turn to the calculation of the ISB transition energies and oscillator strengths. In the top panels of Fig. 3, the solid curves represent the calculated energies $E_{01} = E_1 - E_0$ and $E_{02} = E_2 - E_0$ as a function of w_L for uncoupled [Fig. 3(a)] and coupled [Fig. 3(c)] ACQWs. The position of the related resonances expected in the absorption spectra are instead shown as dotted curves. The energy difference between the absorption resonance and E_{01} , E_{02} stems from the depolarization shift effect induced by the rather high free-electron density resulting from the local dopant concentration $(n = 7 \cdot 10^{17} \text{ cm}^{-3})$ extending over a width of about 10 nm [13, 25, 32]. The oscillator strengths of the ISB transitions f_{01} and f_{02} have been calculated for the uncoupled case [Fig. 3(b)] and for the coupled case [Fig. 3(d)].

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In the uncoupled regime [Figs. 3(a) and 3(b)], only the transition involving the second level of the wide well has a non-zero oscillator strength, because the fundamental state is always localized in the wide well and has no overlap with the first level of the narrow well. This transition corresponds to E_{02} for $w_L < 14$ nm and to E_{01} for $w_L > 14$ nm. As a consequence, the effect of the depolarization shift, which can be seen only in transitions between states with non-negligible spatial overlap, is absent in E_{01} for $w_L < 14$ nm and in E_{02} for $w_L > 14$ nm. In the coupled ACQW [Figs. 3(c) and 3(d)], both transitions are active due to level hybridization and, for $w_L < 14$ nm, they acquire comparable oscillator strengths. A peak value of $f_{02}/f_{01} \sim 0.9$ is predicted at $w_L \approx 10$ nm.



Figure 3: Transition energy and oscillator strength as a function of w_L , calculated (a, b) for $b_t = 10$ nm and $x_{Ge} = 0.81$ and (c, d) for $b_t = 2.3$ nm and $x_{Ge} =$ 0.87. Solid red (green) curve refers to the E_{01} (E_{02}) transition. In panels (a) and (c) the energy of the calculated E_{01} and E_{02} absorption resonances are represented by dotted curves. In panel (d) the ratio between the two oscillator strengths is also reported. The oscillator strength of each transition varies between 0 and $1/m^*$ where $m^* = 0.135$ in electron-mass units is the electron effective mass along the confinement direction, resulting from the values for the L inverse mass tensor reported in Ref. 27.

ISB absorption measurements

The conceptual framework introduced in the previous section is now used to interpret the experimental ISB absorption spectra of Ge/SiGe ACQW samples measured by FTIR spectroscopy at T=10 K. At this temperature excited states are empty, so that only transitions having E₀ as initial level are active. The experimentally determined 2D absorption coefficient spectra $\alpha_{2D}(\omega)$, reported in Figs. 4(a) and 5(a), are fitted with the sum of two Lorentzian peaks (identified with the index *i* = 1, 2), each having, as free parameters, the integrated spectral weight A_i , the half-width-at-half-maximum Γ_i and the peak energy E_{0i}^{abs} . Experimental data for $\alpha_{2D}(\omega)$ were compared with ISBT theoretical absorption spectra obtained with a transition dependent Lorentzian width, set equal to the measured ones (see Fig. 4(b) and 5(b)). To this aim, we note that the $1/\omega$ functional dependence, present in the expression used to model the ISBT absorption coefficient, is filtered out in the experimental spectra since, in this case, $\alpha_{2D}(\omega)$ has been estimated from the dichroic transmittance. The small overestimation of the experimental data in the low energy region of the ISBT spectra can be attributed to this effect.



Figure 4: (a) Dots: experimental ISB absorption spectra measured by FTIR of samples S5 (top), S4 (middle), and S1 (bottom), with Lorentzian fits of the peaks centered at E_{01}^{abs} and E_{02}^{abs} (red and green curves, respectively). (b) Corresponding calculated spectra for w_L = 12 nm and variable thin barrier width b_{t} . (c) Energy difference E_{02}^{abs} - E_{01}^{abs} and (d) ratio of the integrated spectral weight of the two ISB absorption peaks, plotted vs b_{t} . Closed (open) symbols are used for experimental (numerical) data. (e) Calculated squared wavefunctions for sample S5.

In Fig. 4 we analyze ACQWs having the same $w_L = 12$ nm and b_t ranging between 2.3 nm and 4 nm (samples S1, S4, S5). The experimental spectra are very well reproduced by theory. The energy separation $\Delta E^{abs} = E_{02}^{abs} - E_{01}^{abs}$ plotted in Fig. 4(c) decreases for increasing b_t due to the closing of the anticrossing energy gap. Correspondingly, since for $w_L = 12$ nm the E_2 level of the ACQW system has mainly the character of the first excited state in the wide well, we observe an increase of the spectral weight ratio A_2/A_1 (Fig. 4(d)), in agreement with the fact that in the thick-barrier limit, the E_{01} transition is completely suppressed for small values of w_L [see Fig. 3(b)]. Note, however, that $b_i = 4$ nm differs significantly from the uncoupled regime since the spectral weights of the two peaks are still comparable. Indeed, the E_2 and E_1 wavefunctions [see Fig 4(e)] indicate that the excited states at $b_i = 4$ nm are still delocalized in both wells, due to the fact that, for $w_L = 12$ nm, their energy separation in the uncoupled limit is quite small. The absorption-peak linewidths at half maximum Γ_i decrease by increasing the barrier thickness from $\Gamma_i = 7.5$ meV and $\Gamma_2 = 5.5$ meV at $b_t = 2.3$ nm to $\Gamma_i = 4.7$ meV and $\Gamma_2 = 4.1$ meV at $b_t = 4$ nm. We attribute this trend to the reduced impact of IFR scattering on the line-broadening triggered by the decreased value of the electron squared wavefunction at the interfaces (E_2 and E_1 interact with only two interfaces when $b_t \rightarrow \infty$).



Figure 5: (a) Dots: experimental ISB absorption spectra measured by FTIR of samples S3 (top), S1 (middle), and S2 (bottom), with Lorentzian fits of the peaks centered at E_{01}^{abs} and E_{02}^{abs} (red and green curves, respectively). (b) Corresponding calculated spectra at fixed $b_t = 2.3$ nm for different w_L . (c) E_{01}^{abs} (red symbols) and E_{02}^{abs} (green symbols) and (d) ratio of the integrated spectral weight of the two ISB absorption peaks vs w_l . Closed (open) symbols are used for experimental (numerical) data. (e) Calculated squared wavefunctions for sample S2.

In Fig. 5, we study the inter-well coupling as a function of the energy separation between the isolated well levels by comparing the experimental and theoretical absorption spectra of samples with varying w_L and fixed $b_t = 2.3$ nm (samples S3, S1, S2). In reasonable agreement with the theory, we observe in Fig. 5(c) that E_{01} slightly increases with w_L , while E_{02} , involving for $w_{\rm L} < 13$ nm a final state with a character reflecting the one of the first-excited level in the isolated wide well [see Fig. 3(b)], displays a more robust decreasing trend with w_L . The trends reported in Fig. 5(c) result in the decrease of the energy separation of the two absorption peaks, ΔE^{abs} , with increasing w_L , as observed in Figs. 5(a) and 5(b). Correspondingly, when w_L increases from 11.3 to 13 nm, the integrated spectral-weight ratio A_2/A_1 decreases from 0.8 to 0.6 [see Figs. 5(d) and 3(d)], since the level hybridization gets larger and then the $E_2(E_1)$ wavefunction becomes more delocalized, leaking also in the narrow (wide) well, thus reducing (increasing) the $f_{02}(f_{01})$ oscillator strength. Interestingly, the linewidth Γ_2 decreases for increasing w_L , and for $w_L = 13$ nm the peak at E_{02}^{abs} becomes sensibly narrower ($\Gamma_2 = 4.8$ meV) than the peak at E_{01}^{abs} ($\Gamma_{l} = 7.0 \text{ meV}$). This fact can be related to the reduced amplitude of the E_{2} wavefunction at the tunneling barrier interfaces with respect to the E_1 one, as it can be seen in Fig. 5(e). This reduction can be explained considering that the $E_2(E_1)$ level represents the antibonding (bonding) state, resulting from the coupling of the first excited wide well level with the fundamental narrow well level. It follows that the reduced amplitude of the E_2 wavefunction at the tunneling barrier interfaces is due to the increase of the bonding/antibonding interaction when w_L range from 11.3 to 13 nm.

As a final remark, the great accuracy reached by our modeling in terms of *i*) prediction of the absorption energies, *ii*) intensity of the level repulsion, and *iii*) ratio of the oscillator strengths, indicates that our band alignment procedure is quite realistic and, therefore, that our work may be useful in order to definitively establish the *L*-point conduction-band offsets in the SiGe-heterostructure material system. For our samples, we find a discontinuity of 117 meV for $x_{Ge} = 0.81$ and of 80 meV for $x_{Ge} = 0.87$. We emphasize that the growth and characterization of ACQWs is crucial for this purpose, since the ISB absorption spectra of ACQWs are much more sensitive to the precise value of the barrier height, than those measured in isolated wells [33]. The confirmed band offset around 100 meV for *n*-type Ge/SiGe heterostructures makes

them a viable alternative to nonlinear frequency generation of mid-infrared QCLs [34, 35], on the route to a silicon-based THz laser.

Conclusion

We have grown strain-compensated *n*-type Ge/SiGe multi-ACQW samples, and carefully studied, experimentally and numerically, their intersubband absorption spectra, providing a clear interpretative framework of the main experimental features. Our results demonstrate that high-quality tunneling Ge/SiGe heterostructures for electrons at *L*-points of the Ge conduction band can be realized with great reproducibility by CVD. The modeling accuracy in reproducing experimental intersubband absorption spectra guarantees high precision of design and high reliability of material parameters for future SiGe-based THz electronic and photonic devices. In particular the study of wavefunction tunneling in ACQW samples provides reliable values for the estimation of conduction band off-set, which ultimately defines the tunneling barrier height. In summary, our results represent a step forward for the realization of Ge/SiGe THz lasers with broad far-infrared emission range, low refractive index dispersion for frequency-comb operation, and high operating temperature.

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