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Multisubband Plasmons in Doped ZnO Quantum Wells

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Intersubband (ISB) transitions are of high significance for light-emitting and light-detecting devices in the infrared and, when involving large electron densities, for plasmonics and strong light-matter coupling physics. Here it is observed that the simultaneously occurring fundamental and excited-state ISB transitions in highly-doped, m -plane ZnO/Mg_xZn_{1-x}O multiple quantum wells, couple into a single collective oscillation: the multisubband plasmon (MSP). With 2D electron densities up to $4 \times 10^{13} \text{ cm}^{-2}$, an outstanding regime is reached in which the observed MSP frequency is three times larger than that of the fundamental ISB transition as a result of depolarization. This effect is analyzed using a dielectric tensor for ZnO including the interaction of the light with the lattice, the in-plane free electrons, and the off-plane MSP. The impact of the broadening of the MSP and its interaction with phonons is discussed. The results presented here show the potential of ZnO/Mg_xZn_{1-x}O for infrared optoelectronic applications, which can be extended to the THz range with appropriate design of the quantum wells.

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I. INTRODUCTION

Oxides are semiconductors where very high crystallinity control over the heterointerfaces has been achieved recently, allowing phenomena that arise from the presence of a high-mobility two-dimensional electron gas (2DEG) [1,2] as well as from the strong correlation between electrons to be explored [3]. Of particular interest is the ZnO/Mg_xZn_{1-x}O material system, where the quantum Hall effect (QHE) has been observed in heterostructures grown by molecular beam epitaxy (MBE) with electron densities up to $3.7 \times 10^{12} \text{ cm}^{-2}$ [1]. More recently, the observation of the fractional QHE has also been reported in thin layers, where electrons are highly confined and feature mobilities exceeding $1 \times 10^6 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ [3,4], and indeed the rare even-denominator fractional quantum Hall states have been identified [5]. In contrast to the QHE, the fractional QHE can only be explained as a result of the presence of a strong correlation between charge carriers driven by a Coulomb interaction. This strong correlation can be highly

enhanced if, as in the ZnO/Mg_xZn_{1-x}O system, the electrons feature a very large effective mass ($0.24m_0$ in ZnO [6], e.g., much larger than that of GaAs, $0.0637m_0$ [7], where m_0 is the free-electron mass).

The large electron effective mass in the ZnO/Mg_xZn_{1-x}O system is further increased as a result of the high ionicity of these polar oxides, which produces a large coupling between electrons and phonons. This strong interaction can be described by a quasiparticle, the polaron [8], with an effective mass $m^* \approx 0.28 - 0.32m_0$ in ZnO [1,9]. This large electron effective mass implies that the curvature of the subbands in ZnO-based quantum wells (QWs) is so small that the QWs can be heavily doped while keeping the electrons within a narrow energy range. As a consequence, it maintains a large electron population difference between the first and second confined levels in the QW, Δn_{12} . Thus, while Δn_{12} can typically be up to approximately 10^{12} cm^{-2} in GaAs, it can reach values above 10^{13} cm^{-2} in ZnO. The unique combination of large effective mass and very high 2D electron carrier concentrations in ZnO opens a path to explore phenomena that scale with electron density in the QW, such as the depolarization shift in intersubband (ISB) plasmons [10,11], and phenomena occurring in the ultrastrong ISB-phonon coupling regime, i.e., ISB polarons [12].

Here, we explore ISB transitions in Ga-doped, m -plane ZnO/Mg_xZn_{1-x}O multiple quantum wells (MQW)

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featuring very high 2D electron concentrations. ZnO is a material with great potential for ISB transition-based detectors and emitters from the midinfrared to the THz, since it offers several singular characteristics that are not present in the commonly used GaAs-based systems. ISB transition-based absorption and detection have already been observed in ZnO quantum wells [13–15], and the development of quantum cascade lasers in the THz range would benefit from the large phonon energy of ZnO (approximately 72 meV) [16].

The ZnO/Mg_xZn_{1-x}O material system is also chosen in this work because of the availability of nonpolar, native ZnO substrates. To date, ISB transitions in wurtzite wide-band-gap semiconductors, including the GaN and ZnO material systems, have mainly been observed in polar QWs grown along the *c* axis (mostly on foreign substrates, e.g., sapphire or Si), where the quantum confined Stark effect (QCSE) takes place [14,15,17]. Growing the QW structures on a nonpolar orientation (i.e., with the *c* axis parallel to the interface between layers) can suppress the QCSE [18] and should facilitate the ISB transitions due to the absence of internal electric fields that have a detrimental impact on the oscillator strength of the transitions [13]. Growing the QW structures on native substrates should also drastically improve the ISB transition-based devices by reducing the defect density [19] and the residual electron concentration of the films, which is as low as $(1-5) \times 10^{14} \text{ cm}^{-3}$ in the homoepitaxial *m*-ZnO/Mg_xZn_{1-x}O QWs used in this report [20]. The combination of extremely high doping capability of up to $1 \times 10^{21} \text{ cm}^{-3}$ [21] and low residual electron concentration of ZnO and Mg_xZn_{1-x}O provides an unprecedented span of controllable 2D electron concentrations.

II. METHODS

A. Multiple-quantum-well structures

The *m*-plane multiple QW structures presented here, grown by MBE on *m*-plane ZnO substrates, are shown schematically in Fig. 1(a). Samples *A* and *B* feature 3.7-nm-wide QWs and are Ga-doped 1×10^{19} and $2 \times 10^{19} \text{ cm}^{-3}$, respectively. Sample *C* has 4-nm-wide QWs doped $6 \times 10^{19} \text{ cm}^{-3}$ in the central 2 nm of the QW only. An additional undoped sample with nominally identical structure as sample *C* is also used as a reference in the reflectance studies. Finally, sample *D* features 3.9-nm-wide QWs doped $1 \times 10^{20} \text{ cm}^{-3}$.

Figure 1(b) shows an example of the potential profile of one of the QWs presented here as well as the Fermi energy for all samples. The values of ω_{12} and ω_{23} in the QWs are calculated from the self-consistent solution of the Schrödinger and Poisson equations. Both the barrier and QW electron masses are taken to be the ZnO polaron mass $0.28m_0$ [9]. The band offset, ΔE_c , is taken to be $\Delta E_c = 0.675E_g$, since the reported values range from 0.65

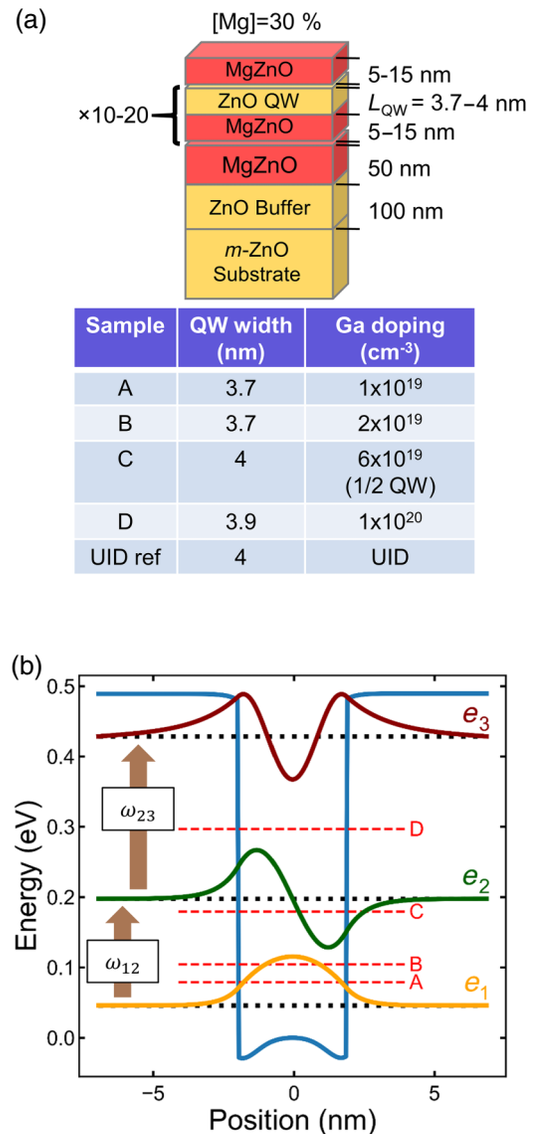


FIG. 1. Details on the samples in this work. (a) Schematic of the multiple quantum well structures presented. The table shows the different doping levels and schemes employed. (b) Potential profile of one of the QWs in sample *D* calculated from the self-consistent solution of the Schrödinger and Poisson equations. Wave functions are offset to match their respective energy levels. Red lines indicate the Fermi energy for all samples. The allowed ISB transitions are also indicated.

to $0.70E_g$ [14,15,22]. The Mg_xZn_{1-x}O band gap is calculated, as in Neumann *et al.* [23], using 25 meV per % of Mg at 30% Mg. With this QW model, the oscillator strength of the e_1 - e_2 transition, f_{12} , is calculated to be between 0.97 and 0.98, and that of the e_2 - e_3 transition, f_{23} , between 1.92 and 1.97. The resulting electronic level structure is similar in all the structures studied here: three conduction-band levels confined in the QWs with an energy difference between the first two, $E_{12} = \hbar\omega_{12}$, ranging from 1220 to

1350 cm⁻¹ (152 to 167 meV) and an energy difference between the second and third levels, $E_{23} = \hbar\omega_{23}$, ranging from 1810 to 1910 cm⁻¹ (224 to 237 meV).

B. Transmittance and reflectance spectroscopy

Slabs 3-mm wide and 6-mm long are cut from the as-grown samples, with the short edges oriented parallel to the c axis. Multipass waveguides are prepared bevelling the short edges at 45° and polishing them mechanically to optical quality with diamond compounds. The back side of the waveguides is polished to optical quality, too. The transmittance is measured at room temperature in a Fourier transform IR (FTIR) spectrometer using a KRS-5, holographic wire-grid polarizer to select the polarization of the incident light. The samples are set in a specially designed blind that allows light only through the bevelled edges of the waveguide.

Reflectance measurements are performed at room temperature on the as-grown samples at 45° and 75° angle of incidence of the light in the same FTIR spectrometer. Since the c axis lies in plane for m -ZnO, in p -polarization transmittance and in all reflectance measurements we place the c axis perpendicular to the electric field [24], such that the material properties in plane and out of plane are isotropic (except those that arise from the ISB transition) and correspond to the $E \perp c$ orientation.

C. Reflectance model

The reflectance spectra are modeled using a semiclassical dielectric function model for each of the constituent layers. For the Mg_xZn_{1-x}O layers, the dielectric function is isotropic (i.e., equal for in plane and off plane of the QW) and includes only the terms accounting for the interaction between the light and the lattice,

$$\varepsilon_{\text{Mg}_x\text{Zn}_{1-x}\text{O}} = \varepsilon_{\infty}^{\text{Mg}_x\text{Zn}_{1-x}\text{O}} \prod_{k=1}^2 \frac{\omega_{\text{LO},k}^2 - \omega^2 - i\gamma_{\text{LO},k}\omega}{\omega_{\text{TO},k}^2 - \omega^2 - i\gamma_{\text{TO},k}\omega}, \quad (1)$$

where $k = 1, 2$ stands for the two phonon modes of nonpolar Mg_xZn_{1-x}O for $E \perp c$, as in Ref. [25]. $\varepsilon_{\infty}^{\text{Mg}_x\text{Zn}_{1-x}\text{O}}$ is the high-frequency dielectric constant, $\omega_{\text{LO},k}$ and $\omega_{\text{TO},k}$ are the LO and TO phonon frequencies of the k mode, and $\gamma_{\text{LO},k}$ and $\gamma_{\text{TO},k}$ are the LO and TO phonon broadenings of the k mode (see [24] for more details on these figures).

In the case of ZnO, the dielectric function is anisotropic, having a different form in the direction parallel to the QW planes, $\varepsilon_{\text{in plane}}$, and perpendicular to the QW planes, ε_z . Both dielectric functions include the background, high-frequency dielectric constant, $\varepsilon_{\infty}^{\text{ZnO}}$, and a harmonic oscillator term analogous to that in Eq. (1) to account for the light-phonon interaction [first term in Eqs. (2a) and (2b)]. However, ε_z features an additional harmonic oscillator term to account for the interaction of the light with the

ISB transitions [second term in Eq. (2a), where the sum in j extends to the first two allowed transitions: $e_1 \rightarrow e_2$ and $e_2 \rightarrow e_3$], and $\varepsilon_{\text{in plane}}$ a Drude model term for the in-plane interaction with the electrons [second term in Eq. (2b)] [12,26,27]:

$$\varepsilon_z(\omega) = \varepsilon_{\infty}^{\text{ZnO}} \frac{\omega_{\text{LO}}^2 - \omega^2 - i\omega\gamma_{\text{LO}}}{\omega_{\text{TO}}^2 - \omega^2 - i\omega\gamma_{\text{TO}}} + \sum_j \frac{\omega_{p,j}^2}{\omega_j^2 - \omega^2 - i\omega\gamma_j}, \quad (2a)$$

$$\varepsilon_{\text{in plane}}(\omega) = \varepsilon_{\infty}^{\text{ZnO}} \frac{\omega_{\text{LO}}^2 - \omega^2 - i\omega\gamma_{\text{LO}}}{\omega_{\text{TO}}^2 - \omega^2 - i\omega\gamma_{\text{TO}}} - \frac{\omega_p^2}{\omega^2 + i\omega\gamma_{\text{in plane}}^p}. \quad (2b)$$

Here γ_j is a phenomenological ISB transition broadening factor we take to be equal to the FWHM of the experimental ISB transition as extracted from the p : s polarization absorbance spectra ratio and $\gamma_{\text{in plane}}^p$ stands for the broadening of the plasma resonance in the in-plane direction. ω_p is the plasma frequency of the in-plane light-electron interaction, defined as $\omega_p^2 = n_{3\text{D}}e^2/m^*\varepsilon_0$, with $n_{3\text{D}}$ the three-dimensional electron concentration in the QW, e the electron charge, and ε_0 the permittivity of vacuum.

Some of the samples presented here feature doping levels so high it is expected that the transitions between excited levels in the QWs will have a significant impact. In this case, dipole-dipole Coulomb interaction couples the optically active transitions resulting in the appearance of a unique collective mode: a multisubband plasmon (MSP) [28]. This MSP can be thought of as an ISB transition in a QW with only two (virtual) confined levels separated by a frequency Ω_{ISBT} . The term of the dielectric function corresponding to this mode can be expressed as a single harmonic oscillator as

$$\frac{\Omega_p^2}{\Omega_{\text{MSP}}^2 - \omega^2 - i\omega\Gamma_{\text{MSP}}} = \sum_j \frac{\omega_{p,j}^2}{\omega_j^2 - \omega^2 - i\omega\gamma_j}, \quad (3)$$

with an effective plasma frequency defined as a function of the plasma frequencies of the individual ISB transitions, $\omega_{p,j}$, as

$$\Omega_p^2 = \sum_j \omega_{p,j}^2 = \sum_j \frac{f_j \Delta n_{2\text{D},j} e^2}{m^* L_{\text{eff},j} \varepsilon_0}. \quad (4)$$

Here, $\Delta n_{2\text{D},j}$ is the difference in the two-dimensional electron density between the j and $j + 1$ levels. $L_{\text{eff},j}$ is the effective width over which the ISB transitions take place, defined as

$$L_{\text{eff},j} = \frac{\hbar}{2S_{j,j+1}m^*\omega_j}, \quad (5)$$

where ω_j are the frequencies of the e_j - e_{j+1} ISB transitions,

$$S_{j,j+1} = \int_0^\infty \Psi_j(z) \Psi_{j+1}(z) \int_0^z \int_0^{z'} \Psi_j(z'') \Psi_{j+1}(z'') dz'' dz' dz \quad (6)$$

and $\Psi_j(z)$ and $\Psi_{j+1}(z)$ are the electron wave functions corresponding to the levels involved in the e_j - e_{j+1} transition [10,11]. Ω_{MSP} is in turn the harmonic mean of the ISB transitions weighted by their respective plasma frequencies,

$$\Omega_{\text{MSP}}^2 = \frac{\sum_i \omega_{p,j}^2}{\sum_j \omega_{p,j}^2 / \omega_j^2}. \quad (7)$$

The reflectance spectra are modeled with the dielectric function approach described above in a transfer matrix formalism (see Ref. [29] and references cited therein) taking into account the whole multilayer structure. The substrate is considered as a semiinfinite medium to account for the light dispersion at the unpolished back side. The fact that the MSP takes place in a width $L_{\text{eff}}^{\text{MSP}}$ different from the QW width, L_{QW} , is taken into account in the model by setting the central $L_{\text{eff}}^{\text{MSP}}$ of each QW as the doped material and the rest of the QW as the undoped material. $L_{\text{eff}}^{\text{MSP}}$ is extracted from Eq. (4) defining $\Omega_p^2 = f_{\text{MSP}} \Delta n_{2\text{D}} e^2 / m^* L_{\text{eff}}^{\text{MSP}} \epsilon_0$ by analogy to the individual $\omega_{p,j}$ and making (a) $f_{\text{MSP}} \approx 1$ because the MSP model assumes there are only two (virtual) electronic levels in the QW, and (b) $\Delta n_{2\text{D}} \approx n_{2\text{D}}$, yielding $1/L_{\text{eff}}^{\text{MSP}} = \sum_j (f_j \Delta n_{2\text{D},j} / L_{\text{eff},j}) / n_{2\text{D}}$. Since the MSP treatment reduces to that of a single ISB transition for lowly doped samples, it is used for all samples in this study.

The $\text{Mg}_x\text{Zn}_{1-x}\text{O}$ in the barrier layers, the undoped ZnO in the substrate, and the fraction of the QW outside $L_{\text{eff}}^{\text{MSP}}$ are homogeneous and isotropic [i.e., the same off-plane and in-plane dielectric function, since $\omega_p = \Omega_p = 0$, and Eqs. (2a) and (2b) are then identical]. In contrast, in the doped ZnO in the QW $\omega_p \neq 0$ and $\Omega_p \neq 0$, and it has to be described with both the off-plane [Eq. (2a)] and in-plane [Eq. (2b)] dielectric functions.

Setting $\omega_p = \Omega_p = 0$ for ZnO in the QWs of the undoped reference sample, the frequencies and broadenings of the light-phonon interaction terms of both ZnO and $\text{Mg}_x\text{Zn}_{1-x}\text{O}$ are extracted by fitting the model to the experimental reflectance spectra as shown in Figs. 2(a) and 2(b), yielding values in agreement with the literature [16,25]. The values thus obtained [24] are then used as a reference for the analysis of the doped samples and ω_p is extracted for all the samples presented here by fitting their reflectance spectra (Fig. 3). The resulting $n_{3\text{D}}$ are then used to calculate the individual $\Delta n_{2\text{D},j}$, which were subsequently fed into Eq. (4) to calculate Ω_p . As discussed in

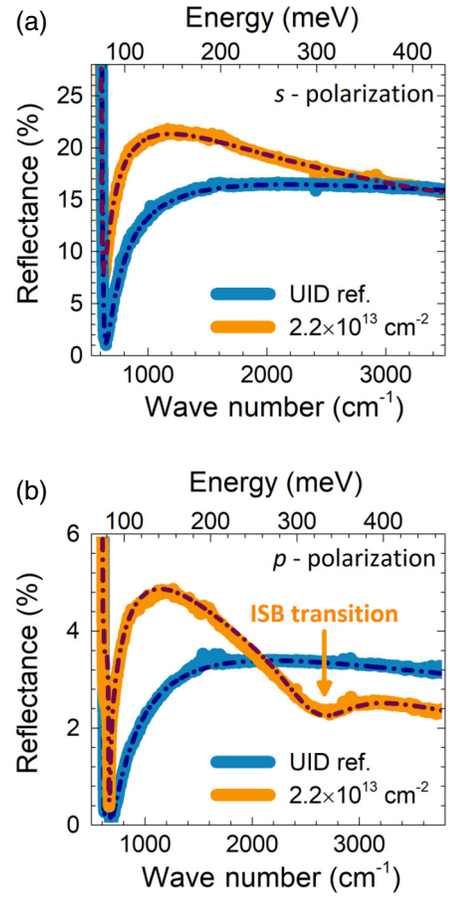


FIG. 2. IR reflectance measurements and fits. (a) Sample C and its equivalent undoped reference under s polarization of the incident light. (b) Same as (a) but under p polarization. The best fit for each spectra is overlaid on the experimental data in all the figures (dashed lines). Legends show the values of $n_{2\text{D}}$ extracted from reflectance experiments in units of cm^{-2} (see Sec. II and Ref. [24]).

[24], the detection limit for the QW 2D electron concentration is approximately $(0.3\text{--}0.6) \times 10^{12} \text{ cm}^{-2}$, and therefore the technique is not sensitive to the residual electron concentration in these films [$(\sim 0.4\text{--}2) \times 10^8 \text{ cm}^{-2}$].

D. Calculation of the dispersion relation from the absorption spectra

The dispersion relation of the MSP is calculated from the simulated absorption spectra, $\alpha(\omega)$, as a function of the 2D electron concentration in the QWs. The absorption spectra are in turn calculated from the off-plane dielectric function of the ZnO QWs [Eq. (2a)] as [29,30]

$$\alpha(\omega) = -\text{Im} \left[\frac{\epsilon_\infty}{\epsilon_z} \right] n_b \frac{\omega \sin^2 \theta}{c \cos \theta}, \quad (8)$$

where n_b is the background, frequency-independent refractive index, c is the speed of light in vacuum, and θ is the

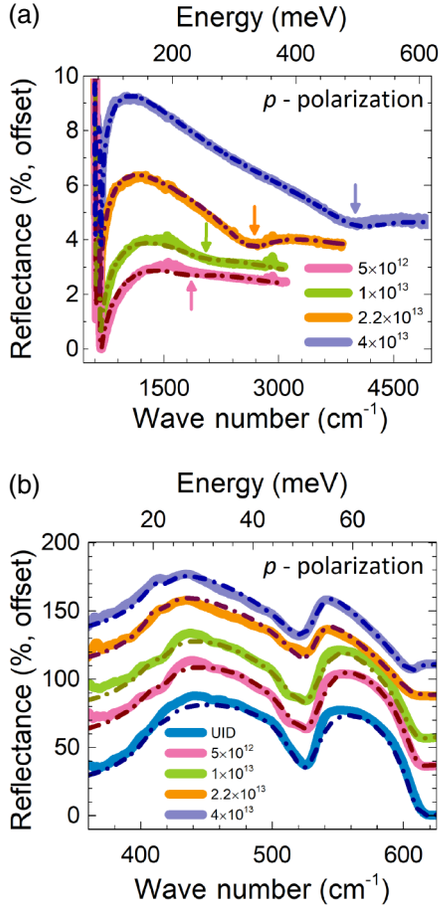


FIG. 3. IR reflectance results. (a) All doped samples in the study under p polarization, offset for clarity. Arrows indicate the presence of the MSP. (b) Enlarged view of the reststrahlen band also under p polarization, offset for clarity. The best fit for each spectra is overlaid on the experimental data in all the figures (dashed lines). Legends show the values of n_{2D} extracted from reflectance experiments in units of cm^{-2} (see Ref. [24]).

angle of incidence of the light with respect to the normal to the plane of the QWs.

III. RESULTS AND DISCUSSION

The room-temperature 45° IR reflectance spectra of the samples presented in this work are shown in Fig. 3(a) for p polarization of the incident light. In all the spectra, the high-energy side of the reststrahlen band is seen at approximately 600 cm^{-1} . The bump at approximately 1500 cm^{-1} is mostly due to the collective interaction of the free electrons with the light along the QW planes. A reflectance dip is observed—especially in the highly doped samples—that is not present in the spectra taken under s polarization, fulfilling the polarization selection rule for

MSP resonance. Its presence in the samples with the lowest 2D electron concentration is also confirmed below with absorption spectroscopy.

Figure 3(b) shows the reflectance spectra at 45° under p polarization across the reststrahlen band. The main features apparent in all the spectra in the figure are also present in the spectrum corresponding to the undoped reference sample and is therefore explained by the interaction of the light with the phonon modes of doped ZnO, undoped ZnO, and $\text{Mg}_x\text{Zn}_{1-x}\text{O}$. Note that the frequencies of interest for this work lie relatively close to the reststrahlen band, which spans from 409 to 590 cm^{-1} in ZnO, i.e., from the $E_1(\text{TO})$ to the $E_1(\text{LO})$ frequencies in the measurement geometry employed here (see Sec. II and [24]). The reflectance valley around 525 cm^{-1} is due to the overlap between the reststrahlen bands corresponding to the two phonon modes in $\text{Mg}_x\text{Zn}_{1-x}\text{O}$ when $\mathbf{E} \perp c$ [25]. The only effect electrons have in these spectra is that of their interaction with the in-plane component of the electric field of the light, resulting in a tilt of the reflectance spectra which, in relative terms, becomes more intense at low-energy end of the reststrahlen band with increasing electron concentration.

Our use of a semiclassical approach with an anisotropic dielectric function to represent the interaction of light with in-plane free electrons and off-plane MSP reproduces very well the reflectance spectra of the doped sample under both s and p polarizations, including the frequency and line-shape of the ISB transitions [Fig. 3(a)]. The obtained values of the plasma frequency, ω_p , and n_{2D} are summarized in [24].

Note that for sample D we obtain $n_{2D} = 4 \times 10^{13} \text{ cm}^{-2}$. Considering a monolayer of m -plane ZnO is approximately 0.21-nm thick, each monolayer of ZnO would then contribute approximately 2×10^{12} electrons cm^{-2} to the total two-dimensional electron density in the QW. This is to be compared to the case of GaAs, where the large two-dimensional electron densities reported in ultrastrong coupling studies yield a contribution that approximately equals 3×10^{11} electrons cm^{-2} per monolayer [31], i.e., roughly one order of magnitude smaller than in ZnO.

Figure 4(a) shows the room-temperature absorbance of sample C in a multipass waveguide configuration for p and s polarizations. In both cases, the absorbance increases for low energies due to absorption by free electrons: note in this configuration the light travels along 6 mm of material and, therefore, the samples are essentially opaque for wave numbers of the light below approximately 1000 cm^{-1} . There is an outstanding feature around 2700 cm^{-1} (335 meV) which only appears in the spectrum measured under p polarization, fulfilling the selection rule for MSP resonances. The MSP resonance can be better visualized if the ratio of the absorbance spectra taken at p and s polarizations is plotted [Fig. 4(b)], which allows measurement of the FWHM and verification that only one absorption peak is present (in agreement with

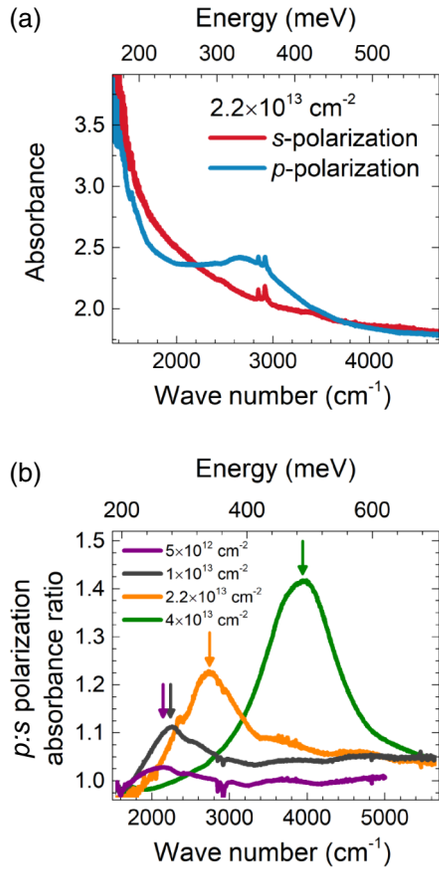


FIG. 4. MSP absorption. (a) Absorbance spectra of sample C under p and s polarization of the incident light in a 45° multipass waveguide configuration. (b) $p:s$ polarization absorbance ratio for all the doped samples in the study. Legends show the experimental values of n_{2D} .

what is expected from a MSP resonance). These peaks shift to higher energies as the 2D electron concentration is increased. Note that the peak energies correspond to what is observed in the reflectance experiments, and also that, consistently with the calculations of the MSP absorption coefficient, the MSP peak height increases with the electron concentration in the QW [32]. The FWHM of the measured MSP lies in the range of 800 cm^{-1} , i.e., from $\gamma_{\text{ISBT}}/\tilde{\omega} \approx 0.38$ in the samples with the lowest n_{2D} to $\gamma_{\text{ISBT}}/\tilde{\omega} \approx 0.28$ in the samples with the largest n_{2D} , where $\tilde{\omega}$ is the observed MSP peak frequency. This is similar to what was obtained by Belmoubarik *et al.* [14] in ISB transition photocurrent experiments at 18 K on c -plane ZnO QWs ($\gamma_{\text{ISBT}} \approx 800 \text{ cm}^{-1}$, $\gamma_{\text{ISBT}}/\tilde{\omega} \approx 0.3$), and lower than the values obtained by Zhao *et al.* [15] in room-temperature absorption experiments, also in c -plane ZnO QWs ($\gamma_{\text{ISBT}} \approx 1250\text{--}1450 \text{ cm}^{-1}$, $\gamma_{\text{ISBT}}/\tilde{\omega} \approx 0.4$).

The ω vs n_{2D} dispersion relation of the MSP transitions, calculated from the absorption spectra of ZnO considering only the off-plane component of the dielectric function [see Sec. II and Eqs. (2a) and (8)], is now

compared to the experimental data. Figure 5 (top) shows the spectral intensity of the MSP resonances together with the experimental results for the MSP peak energy extracted from Figs. 3(a) and 4(b) plotted against n_{2D} . The experimental data agrees excellently with the MSP model, where due to dipole-dipole interaction the oscillator strength is transferred from the lower- to the higher-energy transition in the QW as the Fermi energy moves towards and above e_2 [Fig. 1(b)]. Indeed, in the most heavily doped sample an independent transition model yields two absorption peaks with comparable intensities and at lower energies than the observed peak (see [24]), as opposed to the experiment. Our data show that when the depolarization-shifted independent transitions become resonant, the system becomes transparent for the transitions between adjacent levels, in agreement with the results in Li and Ning [33].

Moreover, the model employed here predicts that the intensity of the observed MSP increases with the 2DEG density, in agreement with the absorption curves in Fig. 4. This is also in agreement with the paper by Załuzny [34] where it is found that for samples where the Fermi energy is above the first excited level of the QW, there are two resonances: one that shifts to higher energies and whose intensity increases with electron concentration, and another one whose energy and intensity decrease with increasing electron concentration. A more recent theory paper by Alpeggiani and Andreani [35] agrees with these observations as well. Also, note the unusual regime reached at these very dense 2DEGs where the depolarization-shifted MSP transition energy is up to three times the energy of the fundamental ISB transition for the most heavily doped sample.

The off-plane component of the dielectric function includes both the terms accounting for the MSP and the interaction of the light with the phonons. The model, therefore, predicts a coupling between these two oscillators. As n_{2D} is increased, the LO-phonon resonance moves across the reststrahlen band towards ω_{TO} as a result of coupling (Fig. 5, bottom). This and the MSP resonances have similar intensities in the range of 2D electron concentrations from 10^{12} to 10^{13} cm^{-2} , i.e., similar to the n_{2D} ranges presented here. However, as a result of the large contribution of the ZnO substrate to the reflectance spectra together with the large broadening of the MSP, the lower branch of the dispersion relation (Fig. 5, bottom) is not observed in these samples. The broadening of the MSP also outshines the impact that the coupling with phonons is predicted to have on its frequency peak: approximately 40-cm^{-1} MSP peak frequency shift vs approximately 800-cm^{-1} MSP broadening. Its observation would thus require a more sophisticated system akin to the ISB polaron laser reported by Liu *et al.* [36], probably the first clear and unambiguous demonstration of an ISB polaron. In their work, a three-level GaAs/(Al,Ga)As intersubband

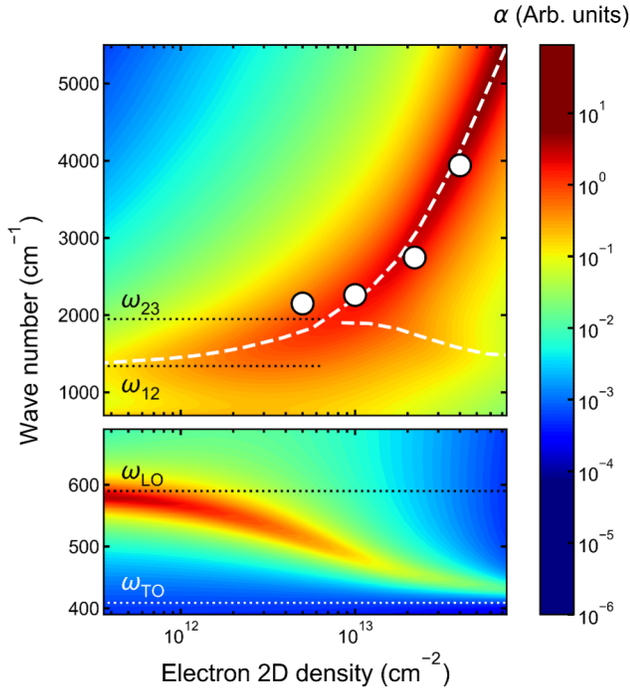


FIG. 5. MSP dispersion relation. ω to n_{2D} dispersion relation calculated from Eq. (8). (Top) MSP resonance calculated expressing ϵ_z as a sum of Lorentzian resonators as in the left-hand side of Eq. (3). The dotted horizontal lines indicate the original energies of the $e_1 \rightarrow e_2$ and $e_2 \rightarrow e_3$ transitions calculated from the self-consistent solution of the Schrödinger and Poisson equations. Dashed white lines indicate the maxima on the absorption spectra. Note how when the MSP reaches ω_{23} the $e_1 \rightarrow e_2$ transition gradually disappears (bottom). Evolution of the phonon resonance with electron density due to the MSP-phonon coupling. Horizontal lines indicate the $E_1(\text{LO})$ and $E_1(\text{TO})$ frequencies from the ZnO QW.

laser was demonstrated at a temperature of 80 K where the Stokes Raman shift (i.e., the difference between the pump and lasing energies) was designed to be around the GaAs LO-phonon energy. It was found that, instead of having lasing action only when the Stokes Raman shift was equal to the LO-phonon resonance, the device would lase when the Stokes Raman shift was equal to the resonances of a dielectric function analogous to the (off-plane) one employed here. It can be foreseen that the same effect would be observed in a hypothetical ZnO/Mg_xZn_{1-x}O three-level laser designed with the Stokes Raman shift around the LO-phonon energy of ZnO. In the ZnO case, though, the ISB polaronic gap for the Stokes Raman shift would be larger than 14 meV (as opposed to approximately 6 meV in the paper by Liu *et al.*) if the transition from the second to the first level of the three-level system is tuned to $\sqrt{\omega_{\text{LO}}^2 - \omega_p^2}$. This difference in the width of the ISB polaronic band gap between GaAs and ZnO would mainly be due to the larger difference between ϵ_∞ and $\epsilon(0)$ in ZnO.

IV. CONCLUSIONS

ISB transitions are demonstrated in m -plane homoepitaxial ZnO/Mg_xZn_{1-x}O MQWs with 2D electron densities in the QWs as high as $4 \times 10^{13} \text{ cm}^{-2}$. It is found that the simultaneously occurring fundamental and excited-state ISB transitions couple into a single resonance, the multisubband plasmon, whose frequency is up to three times that of the fundamental transition. By means of a dielectric function anisotropic with respect to the growth plane of the crystal, the reflectance spectra are analyzed and serve to accurately quantify the actual electron concentrations in the QWs and to determine the ω to n_{2D} dispersion relation of the MSP resonance. The results shown illustrate the potential of m -plane ZnO/Mg_xZn_{1-x}O QWs for optoelectronic applications in the IR, which can be extended to the THz range by an appropriate design of the QW. The implications of these high 2D electron densities and the potential of ZnO to explore ISB-phonon coupling phenomena are discussed.

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