Tailoring Heterovalent Interface Formation with Light

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Integrating different semiconductor materials into an epitaxial device structure offers additional degrees of freedom to select for optimal material properties in each layer. However, interfaces between materials with different valences (i.e. III-V, II-VI and IV semiconductors) can be difficult to form with high quality. Using ZnSe/GaAs as a model system, we explore the use of ultraviolet (UV) illumination during heterovalent interface growth by molecular beam epitaxy as a way to modify the interface properties. We find that UV illumination alters the mixture of chemical bonds at the interface, permitting the formation of Ga-Se bonds that help to passivate the underlying GaAs layer. Illumination also helps to reduce defects in the ZnSe epilayer. These results suggest that moderate UV illumination during growth may be used as a way to improve the optical properties of both the GaAs and ZnSe layers on either side of the interface.

Various semiconductor materials are commonly incorporated into semiconductor devices to meet their performance needs. Material selection is made on the basis of electronic structure and optical and electrical properties, although consideration is typically given to matching crystal structure or thermal expansion coefficients to allow for monolithic growth in a single epitaxial stack. More often than not, these materials are restricted to specific classes of semiconductors, such as group IV, III-V or II-VI semiconductors, for ease of growth. However, there are obvious advantages for incorporating materials from several classes to select for very specific properties in each layer and/or the substrate. This includes the growth of III-V optoelectronic devices on Si substrates and the application of high bandgap II-VI cladding layers on III-V absorbers or light-emitting active layers. Combining dissimilar materials therefore creates myriad possibilities for designing next generation semiconductor devices if the integration can be carried out successfully.

Interfaces are an inevitable product of material integration, and the formation of heterovalent interfaces presents a particular difficulty for growth. Polar heterovalent interfaces composed of only one bond type will result in large charge imbalances, so they ideally will be formed with some intermixing over a few monolayers. Achieving the optimal distribution of bonds across the interface while minimizing defect formation requires precise control of the interface initiation conditions. For example, point defect and secondary phase formation as well as atomic diffusion between layers can occur during the growth of a ZnSe/GaAs hetero-structure by molecular beam epitaxy (MBE) if the interface is not initiated properly. Such defects would be detrimental to device performance if the heterovalent interface is located between critical layers.

Two free parameters that are typically available during growth to provide this control are temperature and molecular or atomic flux. Temperature is the easiest to adjust and strongly influences most growth processes. High growth temperatures are advantageous for enhancing adatom surface mobility and producing high quality crystalline epilayers. Yet, elevated temperatures can also promote atomic inter-diffusion between adjacent layers, which can degrade their optical and transport properties. It may therefore be beneficial to use lower growth temperatures to achieve abrupt interfaces. Low temperatures can additionally lead to more abrupt doping profiles and extend the critical thickness of lattice mismatched layers, but they can also inhibit adatom surface migration, promote three-dimensional growth and produce higher defect concentrations. The elemental source flux ratio may also be adjusted to modify surface migration, defect formation and the bonding environment at the interface, although care must be taken to avoid unwanted micro- and nano-structure formation within the epilayer. Optimal control of growth processes therefore may not be achieved by varying these two parameters alone.

Light stimulation of the growth surface offers an additional parameter with which to alter growth processes. In particular, it has been shown to improve the crystal quality of II-VI semiconductor epilayers grown at temperatures as low as 150 °C and has helped to improve extrinsic doping efficiency. This approach appears to be advantageous for forming heterovalent interfaces because it permits the use of low growth temperatures.
without sacrificing material quality. Nevertheless, most research has focused on the effect of light stimulation on the II-VI epilayer only, and very little attention has been paid to the effect of light on the initiation of a II-VI/III-V interface or its affect on the underlying III-V material. Additional information about its potential for modifying interface formation is needed in order to consider it as a tool for aiding material integration.

Herein, we present an investigation into the mechanisms by which light stimulation modifies low temperature growth of II-VI/III-V interfaces, using ZnSe/GaAs as a model system. Its effects on the optical, structural and morphological properties of both epilayers as well as the interface were studied as a function of illumination conditions. We find that UV illumination alters the starting coverage of the As-reconstructed GaAs surface at the start of the initiation process, leading to a different mixture of bonds at the interface. This results in an apparent improvement in the passivation of the underlying GaAs epilayer. UV illumination also improved the optical properties of the ZnSe epilayer. The combination of low-temperature growth and light stimulation therefore provides a way to control the interface formation and achieve high material quality. Based on these findings, we offer some guidance for using light-stimulated growth techniques for heterovalent interface formation.

Results and Discussion

ZnSe epilayers were grown by MBE on GaAs epilayer/GaAs substrate structures. Initiation of the ZnSe/GaAs interface commenced with a Zn pre-exposure of an As-terminated GaAs surface, followed by growth of the ZnSe layer. Details of the growth are included in the Methods section and are illustrated in Figure 1. A Xe lamp was used as the illumination source, and the illumination conditions during the interface initiation process were varied to determine the mechanisms by which UV illumination modifies its properties. We investigated three scenarios: (1) no light exposure (referred to as “dark reference”), (2) light exposure starting after the Zn pre-treatment...
(referred to as “dark-start”) and (3) light exposure starting before the Zn pre-treatment phase (referred to as “light-start”).

Structural Properties. Figure 2(a,b) shows ω-2θ high-resolution X-ray diffraction (HRXRD) curves of light-start and dark-start samples, respectively. The thicknesses of the ZnSe epilayers were the same regardless of the starting condition or the lamp power. The primary difference between the two starting conditions is the appearance of weaker Pendellösung fringes in the HRXRD curves of the light-start samples, indicating diffusion at the interface. The fringes remained clear in all dark-start samples, indicating the presence of an abrupt interface. (c,d) Tilt-view SEM images of the light-start sample grown under a lamp power of 150 W as well as a dark reference sample. The yellow curves display a first derivative of the image contrast profile at the ZnSe/GaAs interface. The contrast profile is sharp at the interface of the dark reference sample.
sample but was graded at the interface of the light-start sample. Together, these results indicate that the degradation of the Pendellösung fringes in the HRXRD curves is due to compositional grading at the ZnSe/GaAs interface. Higher lamp power and light-start conditions both appear to enhance the grading effect.

The growth temperature used here (200 °C) is far below the optimal ZnSe growth temperature (300 °C) and, in general, lower growth temperatures are advantageous for creating abrupt ZnSe/GaAs interfaces. Thus, the role of illumination before and during the Zn pre-treatment phase is somewhat similar to increasing the substrate temperature, which promotes more interface intermixing. Excessive atomic intermixing at the interface was also observed by our group when growing ZnSe epilayers on GaAs surfaces that had been depleted of As coverage through thermal annealing at temperatures ≥500 °C. In ref. 2, strong Pendellösung fringes were observed when ZnSe epilayer growth was initiated on a fully As-reconstructed surface with a Zn exposure at 300 °C. The predominant mixture of As and Zn atoms suggested that a thin layer (on the order of a monolayer) of Zn₃As₂ or ZnASₓ may develop at that step, aiding abrupt interface formation. However, the width of the ZnSe HRXRD peaks broadened and the Pendellösung fringes between the ZnSe and GaAs peaks disappeared when the As coverage was reduced through annealing at elevated temperatures (500 °C and 670 °C). That evolution was attributed to depletion of the As coverage at the start of the interface initiation, which would allow more intermixing of As, Ga, Zn and Se. Although interface initiation is carried out at even lower temperatures in the present case, illumination during the Zn pre-treatment step can also affect the As coverage. UV illumination is known to selectively enhance anion desorption from the surface via disruption of the anion bonds by photo-generated holes and has been observed specifically to promote As desorption from GaAs surfaces by our group. The HRXRD results therefore suggest that UV photon irradiation acts to deplete the As coverage of the reconstructed GaAs starting surface to produce a similar effect at elevated substrate temperatures. On the other hand, the Pendellösung fringes remained strong in the HRXRD curves of all dark-start samples, regardless of the lamp power. This result indicates that a Zn sub-monolayer deposited on a fully As-reconstructed GaAs surface stabilizes the interface against intermixing, and subsequent exposure to light does not substantially affect it.

**Morphological Properties.** Figure 3 shows atomic force microscopy (AFM) images of the light-start samples and their corresponding power spectral densities (PSD). The PSDs provide information about the frequencies of spatial feature wavelengths. The general form of the PSD is derived from continuum growth models and is
expressed as $\delta h \propto \nabla^2 h + \nabla^4 h + \cdots$. To produce PSD curves in a wide range of spatial frequencies, AFM images were obtained over several areas: $1 \times 1 \mu m^2$, $5 \times 5 \mu m^2$ (not shown here), and $10 \times 10 \mu m^2$. Care was taken not to include features from the SeO$_2$ clusters in the analysis. As shown in Figure 3(a–d), the PSD curve slope of the dark reference sample falls between $-2$ and $-4$. This slope moves closer to $-2$ with increasing lamp power as shown in Figure 3, indicating that the surface features become somewhat larger. There is an additional feature in the PSD spectra, marked with boxes in Figure 3(a–d), that represents the periodicity of hillocks in the range of $0.1$–$1 \mu m$ ($1$–$10 \mu m^{-1}$). In the dark reference sample, the two regions of elevated intensity represent the prevalence of two surface hillocks having periodicities of $0.5 \mu m$ ($2 \mu m^{-1}$) and $0.2 \mu m$ ($5 \mu m^{-1}$). An increase in the light intensity broadens and lowers the range of dominant spatial frequencies, indicating that the surface hillocks have generally become wider (i.e. there is a shift to lower spatial frequencies). We should note that the actual surface features associated with the ZnSe epilayer are never more than a few nanometers in height.

Substrate temperature, molecular interactions and elemental surface coverage (i.e. As, Se, etc.) all strongly affect the growth mode. Widening of the surface features with increasing lamp power could be due to one of two factors: enhanced dissociation of multiatomic Se molecules or improvement in adatom mobility. Regarding the first possibility, the dissociation energy of multiatomic Se molecules is $3.55$ eV$^{27}$. Thus, direct absorption of photons with energies greater than $3.55$ eV may lead to higher cracking rates. However, the flux of photons in this energy range is limited in the Xe lamp emission spectrum. Furthermore, growth was carried out with a standard valved-cracker cell held at $1000$ °C, which helps to improve dissociation of multiatomic Se molecules in the first place. Thus, this scenario seems unlikely to be the primary pathway for growth mode modification.

Instead, the growth mode may be modified by improving the mobility of surface adatoms. Low temperatures tend to reduce adatom mobility, leading to 3D growth modes and roughening. This is the case in the present situation, where the ZnSe epilayers were grown well below the optimal growth temperature ($200$ °C instead of $300$ °C). Standard growth of ZnSe also relies on a high Se/Zn flux ratio, which leaves the surface Se terminated. While UV illumination enhances anion desorption, the cation desorption rate is less affected$^{28,29}$. The resulting surface becomes more metal-rich, leading to improved adatom mobility and migration and to smoother surfaces$^{30}$. This appears to be a plausible explanation for the broadening of the surface features. Figure 4 shows surface AFM

Figure 4. The left-hand portion of the figure displays $1 \times 1 \mu m^2$ and $10 \times 10 \mu m^2$ surface AFM images of the dark-start samples. (a–d) Displays their corresponding PSD curves. The curve slopes approached 2 with increasing Xe lamp power. Regions of elevated PSD shifted to lower spatial frequency with increasing lamp power, which indicates that the period of the hillocks on the sample surface increased. (e) The PSD slopes of the samples as a function of Xe lamp power. The PSD curve slopes of the light-start and dark-start samples decreased with increasing Xe lamp power.
images of the dark-start samples and their corresponding PSD curves. The behavior is much the same as that of
the light-start samples. For a given lamp power, there is a slightly greater decrease in the PSD slope compared to
the light-start samples, shown in Figure 4(e), indicating a smoother surface with wider hillock structures. This
could be due to a decrease in any roughening that can occur when the interface itself is grown under UV illu-
mination. However, the effect appears to be minimal compared to the overall change generated by illumination
during ZnSe epilayer growth.

Figure 5. (a–f) PL emission from the ZnSe epilayers grown under light-start and dark-start conditions,
respectively. The PL spectra consist of emission from NBE and DLE regions. The DLE region is divided into SA
and Cu-green emission. (g) The NBE/DLE peak intensity ratio of light-start samples as a function of Xe lamp
power. (h) The normalized PL spectrum of dark reference samples grown at 200 °C and 300 °C. With increasing
Xe lamp power, the PL spectrum of the 200 °C-grown ZnSe epilayer approaches that of the 300 °C-grown ZnSe
epilayer, with suppressed I₁deep and DLE emission. (i) DX/I₁deep ratio of light-start and dark-start samples. DX/
I₁deep increase with lamp power approaching that of 300 °C-grown ZnSe.
Optical Properties. Figure 5(a–f) shows the photoluminescence (PL) spectra of the ZnSe epilayers in the light-start and dark-start samples. The emission spectra can be divided into two important regions: near-band-edge emission (NBE) around 2.8 eV and deep-level emission (DLE) below 2.6 eV. Within the DLE region, emission around 1.95 eV is usually referred to as the self-activated (SA) region and has been linked to distant donor-acceptor pair transitions, especially those involving zinc vacancies (V\text{Zn}) and group III donor impurities\textsuperscript{31–33}. The NBE region consists of free and donor-bound exciton emission (FX and DX) peaks as well as deep acceptor-bound exciton (I\text{1}\text{deep}) emission and its phonon replicas. The overall trends in the light-start and dark-start samples are similar. The total emission intensity increases when grown under UV irradiation compared to the dark reference sample. It is generally attributed to a reduction in non-radiative recombination.

We start by inspecting the DLE region. Evolution of the SA emission as a function of lamp power is most evident. It first increases upon irradiation with a lamp power of 80 W and then subsequently decreases with increasing lamp power up to 150 W. It is unclear whether the slight increase at 80 W is due to an increase in the V\text{Zn} or group III (possibly Ga) impurity concentration or to a change in the transfer rate of carriers to those states (as opposed to non-radiative states), which could enhance the emission probability without changing the defect density. The ratios of the intensities of the peaks in the NBE and DLE regions are plotted in Figure 5(g) for both sets of samples. Generally, the NBE/DLE emission does not change significantly as the lamp power is increased from 0 W to 80 W, which suggests that most of the SA emission increase is due to an overall increase in the number of carriers that reach those acceptor and donor states, rather than the density of states themselves. What is clear is that the NBE/DLE ratio substantially increases as the lamp power is further increased to 150 W. That increase is largely caused by a reduction in the SA emission rather than an increase in the NBE. As noted above, excess Se desorption is known to increase under UV irradiation, leading to the possibility of forming a more metal-rich growth surface\textsuperscript{30}. A greater amount of Zn on the growth surface would indeed reduce the concentration of V\text{Zn} in addition to promoting smoother surfaces. To check this analysis, we compared the evolution of the SA emission to another instance where Se desorption and adatom mobility is enhanced: elevated substrate temperatures. The PL spectra of two samples grown under dark conditions at 200 °C and 300 °C are shown in Figure 5(h). The sample grown at 300 °C also exhibits a reduction in the SA emission compared to the dark reference sample grown at 200 °C. A similar trend is observed in the I\text{1}\text{deep} intensity (see Figure 5(i)), which is also associated with Zn vacancies\textsuperscript{34}.

Strong UV photon irradiation also leads to an increase in the DX and FX emission in the NBE region. In particular, the emission shifts from a dominant I\text{1}\text{deep} peak to dominant DX and FX peaks. This is correlated with a decrease in DLE and an overall improvement in material quality. The NBE/DLE ratio, which can serve as a proxy for ZnSe epilayer quality, becomes comparable to that of ZnSe grown in the dark at 300 °C. This is approximately a factor of 4 times the ratio of the dark reference sample.

We note here that the effect of UV irradiation on samples grown under light-start or dark-start conditions are similar, meaning that the improvement is associated with the bulk ZnSe epilayer. The slightly higher NBE/DLE ratio in the dark-start sample grown under a lamp power of 150 W relative to the light-start sample may be due to differences in bonding at the interface. However, there is not enough evidence to suggest this is a statistically significant difference between light-start and dark-start conditions.

Further information about the effect of photon irradiation on the ZnSe/GaAs interface can be obtained from PL measurements of the underlying GaAs, shown in Figure 6. The PL spectra of light-start and dark-start samples as a function of illumination conditions are shown in Figure 6(a–f), and they exhibit three features: DX and FX emission near 1.51 eV, carbon-bound (C\text{as}) emission at 1.49 eV, and a broad emission peak at 1.48 eV that exhibits a low energy tail. The emission peak at 1.48 eV is related to Ga-Se bonding at the interface, while the low energy tail is attributed to a more complex mixture of elemental species\textsuperscript{35}. Both evolve with illumination conditions\textsuperscript{36}. In particular, the intensity of this peak increases substantially when interface initiation is carried out with a light-start at the highest lamp power. The HRXRD measurements show evidence of more elemental intermixing at the interface caused by excessive As desorption under these starting conditions, which could account for additional Ga-Se bond formation. To confirm the origin of this behavior, we simulated conditions of excess As desorption by increasing the As desorption temperature prior to interface initiation. Figure 5(g) shows the evolution of the PL peak at 1.48 eV when the As desorption process was carried out at elevated temperatures up to 670 °C, and the ZnSe epilayer was subsequently grown at 200 °C under dark conditions. This peak becomes evident at desorption temperatures of 500 °C and overwhelms the C\text{as} peak when As desorption is carried out at 670 °C. The intensity of the 1.48 eV peak in the light-start/150 W sample falls somewhere in between these two samples, strongly supporting our claim that the 1.48 eV emission is enhanced by additional As desorption.

Besides changing the amount of As coverage on the starting GaAs surface prior to interface formation, elevated ZnSe growth temperatures can also promote interface intermixing, leading to similar results. The PL spectra of two ZnSe epilayers grown under dark conditions at 200 °C and 300 °C are shown in Figure 6(b). The sample grown at 300 °C also exhibits a moderate PL peak at 1.48 eV, suggesting that Ga-Se bonds have formed across the interface at higher concentrations than in the sample grown at 200 °C, likely due to enhanced diffusion across the interface during ZnSe growth at elevated temperatures.

Comparatively, the dark-start samples do not show such an enhancement in the 1.48 eV peak because the dark-start protects against As-desorption. In Figure 6(d), this portion of the spectrum has been multiplied to show that there is a weak peak near 1.48 eV, but its overall intensity is much lower than the light-start sample grown under similar conditions. This behavior arises through the formation of an abrupt interface upon Zn pre-treatment of an As-rich surface, preventing excessive Ga-Se bonding.

The DX and FX emission of the light-start samples is also enhanced with increasing UV lamp power. This effect is likely the result of lower interface recombination, possibly by Se passivation\textsuperscript{37} or a stable interface formed through Ga-Se bonds\textsuperscript{2, 36} followed by a high quality ZnSe layer. Based on this result, light stimulated growth of the ZnSe epilayer at 200 °C and a light-start appears to help enhance the underlying GaAs epilayer emission. This is especially evident when comparing the ratio of the intensity of the DX and FX excitonic emission to the Ga-Se
peak intensity at 1.48 eV, as shown in Figure 6(i). By contrast, the excitonic emission of the dark-start samples is comparable to or lower than that of the dark reference sample, with only minor changes in the Ga-Se peak intensity. It is possible that a greater percentage of Zn-As bonds increases interface recombination in the dark-start case. In any event, the excitonic/Ga-Se peak intensity ratio of both sample sets is much higher than the dark reference sample grown at 300 °C until the highest lamp intensities are used (also depicted in Figure 6(i)).
It is also noteworthy that C\textsubscript{As} acceptor bound excitonic emission is largely suppressed in the dark-start samples, which it is present in both the dark reference at light-start samples. This effect is highly reproducible, and we have observed it in many additional sample sets not detailed here. Secondary ion mass spectroscopy (SIMS) measurements, shown in Figure 7, indicate that there is no significant difference in the C concentration at the ZnSe/GaAs interface, or within the ZnSe epilayer and GaAs epilayer. We do not understand the origin of this effect, but we currently suspect that it is due to some mechanism that alters exciton binding to C acceptors.

### Figure 7.
Cross-sectional carbon profile of light-start, dark-start and dark reference samples obtained by SIMS. For the light-stimulated samples, the Xe lamp power was 150 W. There is no significant difference among the samples in cross-sectional carbon ion counts.

### Figure 8.
(a–d) Schematic of the light-start and dark-start growth process, respectively. During the light-start process, a Se-induced secondary phase may be formed. Meanwhile, formation of a secondary phase can be inhibited by a sub-monolayer-thick Zn layer.

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### Summary of Results and Discussion.
When considered together, the structural, morphological and optical results presented here suggest that UV photon irradiation modifies the low temperature growth of ZnSe on GaAs through two mechanisms. (1) Optical injection of non-equilibrium concentrations of photogenerated holes drives As desorption from the reconstructed GaAs surface, exposing underlying Ga and promoting a greater distribution of chemical bonds at the interface. This mechanism was verified through evidence of intermixing from HRXRD measurements and increased signatures of Ga-Se bonding in the PL spectra of the GaAs epilayer. UV photon irradiation is also known to disproportionately elevate anion desorption from semiconductor growth surfaces\textsuperscript{25}. This process is depicted in Figure 8. (2) UV photon irradiation during ZnSe deposition improves the material quality, substantially reducing deep level emission in these low temperature-grown epilayers. Improvement in the optical and transport properties has generally been
observed in II-VI semiconductors grown under photon irradiation and is attributed to metal-rich surface conditions created by enhanced anion desorption\(^2\).

The changes observed here in the structural and optical properties of the ZnSe and GaAs epilayers and the interface between them are subtle but important. Photon irradiation allowed the ZnSe epilayer to be carried out at 200 °C rather than 300 °C with little-to-no loss in optical quality. The lower growth temperature then prevented diffusion-driven elemental intermixing at the ZnSe/GaAs interface during growth of the ZnSe epilayer, which introduces a high density of bound defect states in the underlying GaAs, as clearly shown in Figure 6(i). Furthermore, moderate lamp power (for example 115 W in this study, as shown in Figure 6(b)) may be somewhat beneficial for promoting a measured degree of intermixing, reducing recombination at the interface and improving excitonic emission within the GaAs layer. Photon irradiation during growth therefore can be used as a tool to decouple material and interface quality from substrate temperature. In the case of achieving balanced As coverage on the starting GaAs surface to allow some Se passivation while preventing excessive intermixing, it is possible to promote As desorption through elevated temperatures or photon irradiation. However, this would require very precise control of the temperature and time of the desorption process to achieve a favorable As coverage, as demonstrated in our previous work\(^2\). Moderate light intensities, on the other hand, appear to provide some amount of regulation, allowing control over other growth parameters to be relaxed. One could also apply different light levels to the growth of the interface and growth of the ZnSe epilayer to control each separately. The PL spectra from the GaAs epilayers suggests that there is still room for further enhancement of the excitonic emission by precise control of the UV lamp power to balance the bonding environment at the ZnSe/GaAs interface. Overall, our results suggest that photon irradiation can be combined with low temperature growth in a number of ways to tailor the properties of ZnSe/GaAs heterostructures. For example, light can be applied at the interface initiation step to improve photo-passivation of the underlying GaAs epilayer and to improve excitonic properties. This approach may be best for cases where preserving the optical emission efficiency in the III-V layer is of paramount importance (i.e. in applications where the II-VI layer is used to clad the III-V layer). Alternatively, a dark-start interface initiation approach can be used to produce a much more abrupt ZnSe/GaAs interface and suppress C\(_{\text{gax}}\) recombination in the GaAs. Thus, a dark-start may be advantageous from this perspective. More broadly, irradiation of the growth surface with above-bandgap photons may be a useful tool for modifying the structure or chemical composition of a variety of interfaces, which could be increasingly valuable as material integration continues to rise. The new understanding of how light modifies heterovalent interface formation presented here will help to guide the determination of whether photo-assisted growth techniques can provide some benefit to other material systems.

**Conclusion**

In summary, ZnSe epilayers were grown at 200 °C on As-terminated and Zn pre-treated GaAs surfaces under an array of UV photon irradiation conditions. HRXRD, SEM, AFM, SIMS, and PL measurements reveal that photon irradiation during interface initiation (light-start) enhances As desorption and allows additional Ga-Se bonding to occur at the ZnSe/GaAs interface. This approach led to some improvement in the emission from the GaAs epilayer. Under both light-start and dark-start conditions, light stimulation substantially improved the ZnSe epilayer material. These results suggest that low temperature growth accompanied by moderate UV photon irradiation can be used to tailor interface formation and improve the properties of the optical quality of the ZnSe and/or GaAs epilayers.

**Methods**

**Sample Growth.** Samples were grown on semi-insulating (100) GaAs substrates in an Omicron EVO25 MBE system with separate III-V and II-VI semiconductor growth chambers. The two chambers were equipped with conventional K-cells for group-III and/or group-II elemental sources and valved cracker effusion cells for the group-V and group-VI elemental sources. The substrate temperature was measured with a k-Space BandiT band-edge thermometry system with an accuracy of ±1.0 °C. The substrates were first outgassed at 300 °C in an ultrahigh vacuum chamber before being transferred to the III-V growth chamber, where the oxide layer was thermally desorbed at 610 °C for 10 minutes under an As\(_2\) overpressure. A 500 nm-thick homoepitaxial GaAs epilayer was then grown, followed by growth of a thick amorphous As layer at room temperature for one hour which was used to protect the GaAs surface while the sample was transferred to the II-VI growth chamber. Inside the II-VI chamber, the amorphous As cap was desorbed at 330 °C in the absence of any group II or VI overpressure to leave a partially As-terminated surface\(^1\). In a few specified cases, the As cap desorption temperature was varied. Immediately after As desorption, the substrate temperature was set to 200 °C. Interface initiation commenced with exposure of the As-terminated surface to a Zn flux for 2 minutes to form a sub-monolayer (ML) of Zn. This was followed by a 30 seconds exposure to Se, after which the Zn flux was turned back on to begin the growth of a thick ZnSe epilayer. All ZnSe growths were carried out at 200 °C for 30 minutes under Se-rich condition\(^3\), except for one particular case, where the growth was carried out at 300 °C. The resulting epilayers were nominally 230 nm thick. Light was sourced from a Xe lamp (Oriel 6255) operated at three different powers: 80 W, 115 W and 150 W. A 0.6 neutral density (ND) filter was placed between the lamp and the sample, and the beam was partially collimated to a spot size of approximately 25 cm\(^2\) on the sample. The light flux was held constant throughout the duration of the growth.

**Characterization.** The layer thickness was confirmed by SEM image and ω-20 HRXRD curve simulation. The estimated thicknesses of all ZnSe layers were approximately the same. AFM was used to measure the surface morphology of the ZnSe epilayers as a function of the Xe lamp power. Low temperature PL of the ZnSe epilayers was measured with a GaN diode laser (405 nm) operated at an excitation power ~1 mW and using a 435 nm long-pass filter to block the laser line. Photoluminescence of the underlying GaAs epilayers and ZnSe/GaAs interfaces were obtained with a Nd:YVO laser (532 nm) to avoid pumping the ZnSe epilayer. The excitation power was ~1.2 mW, using a 570 nm long-pass filter. Both measurements were carried out with a 150 I/mm grating.
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Author Contributions
K.P. conducted MBE growth and measurements. K.A. and K.P. designed experiments and analyzed measurement results. All authors reviewed the manuscript.

Additional Information
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