Mapping the Origins of Luminescence in ZnO Nanowires by STEM-CL

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Supporting Information

ABSTRACT: In semiconductor nanowires, understanding both the sources of luminescence (excitonic recombination, defects, etc.) and the distribution of luminescent centers (be they uniformly distributed, or concentrated at structural defects or at the surface) is important for synthesis and applications. We develop scanning transmission electron microscopy–cathodoluminescence (STEM-CL) measurements, allowing the structure and cathodoluminescence (CL) of single ZnO nanowires to be mapped at high resolution. Using a CL pixel resolution of 10 nm, variations of the CL spectra within such nanowires in the direction perpendicular to the nanowire growth axis are identified for the first time. By comparing the local CL spectra with the bulk photoluminescence spectra, the CL spectral features are assigned to internal and surface defect structures. Hyperspectral CL maps are deconvolved to enable characteristic spectral features to be spatially correlated with structural features within single nanowires. We have used these maps to show that the spatial distribution of these defects correlates well with regions that show an increased rate of nonradiative transitions.

Zinc oxide (ZnO) offers a number of attractive features relevant to application, including a wide band gap, optical transparency, large exciton binding energy, and earth-abundant elemental composition.1 ZnO nanowires, in particular, have shown promise in optoelectronics,2 lasers,3 photovoltaics,4 and gas sensing.5 However, defects can dictate nanowire device performance: surface states cause band bending and deplete the carrier density,6,7 whereas optically active defects provide unwanted recombination routes. The density and distribution of these defects, as well as the length scale over which they influence properties, are important practical considerations both for applications where surface interactions are critical (e.g., photocatalysis and chemical sensing8–11) and more generally for ZnO nanodevices.

Extensive work on ZnO photo- and cathodo-luminescence (PL and CL respectively) has identified a number of spectral contributions.12 Primary luminescence signals arise from transitions across the band gap often mediated by excitons,12 and transitions are associated with a variety of defects.13 Impurity elements often play a crucial role, but even in elementally pure ZnO there are defect states that lie within the band gap and contribute to luminescence, include those arising at 2.0 eV (doubly charged O vacancies)14 and 1.94 eV (O-rich growth conditions or O interstitials).15,16 In nanostructures, the external surface becomes particularly significant and may be associated with different types (or concentrations) of defects compared with the core. Broad signals from such surface states in nanostructures have previously been observed by PL (at 2.4 to 2.54 eV17–19 and 2.2 eV20 central energies) and monochromatic CL (2.5 eV) in ZnO nanostructures. Despite progress, the assignments of spectral features to specific structures are often uncertain, and a combined understanding of both the spatial distribution and the spectral variation of defects remains elusive.

An ideal technique for characterizing defect states in nanostructures would combine spatially resolved emission measurements with high-resolution structural imaging. Here, we describe scanning transmission electron microscopy–cathodoluminescence (STEM-CL) measurements of individual high-quality ZnO nanowires. Existing efforts to study ZnO luminescence using SEM-CL25,26 suffer from large electron-beam interaction volumes, which limit resolution; CL

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emission has only been mapped longitudinally along single ZnO nanowires. The characteristics in directions normal to the growth axis have not been resolved, although they are critical for both a fundamental understanding and a number of device applications. STEM-CL has not been widely applied to such systems, although it has previously been used to map CL from quantum wells in GaN nanowires and collect localized CL spectra from ZnO tetrapods and 1D hyperspectral line scans on single GeS nanosheets.28

Here, by using STEM-CL, we have deconvoluted the spatial variation of the CL spectra arising from the nanowire core, the nanowire surface, and internal defects. The high-energy electron probe used in STEM affords simultaneous structural imaging with local excitation of electron–hole pairs. The spectrum of light emitted by radiative recombination is recorded as the electron beam rasters the nanowire. The resulting high-resolution maps correlate the CL spectra with the nanowire geometry and hence help to identify the origin of optical traps in individual nanowires.

Samples were prepared as described in the Supporting Information (SI). STEM-CL data were acquired with a Gatan Vulcan CL system in a JEOL 2100 field-emission scanning TEM operated at 80 kV with a 1 nA beam current and a 1 nm spot size. A pair of mirrors mounted on the sample holder focused the emitted light onto optical fibers that fed an optical spectrometer. CL spectra were recorded, one at each pixel rastered by the electron beam (10 nm pixel size, 3 s dwell time resulting in an electron dose of 30 μC/μm²), with 16 × 16 subpixel scanning enabled. These imaging conditions were chosen to maximize the CL signal and spatial resolution while minimizing damage to the nanowire structure. To enhance radiative recombination and minimize knock-on damage, the nanowires were liquid-nitrogen-cooled to a temperature of 103 K. PL was performed at room temperature using a He/Cd laser with an excitation wavelength of 325 nm using a Renishaw fluorescence microscope.

High-quality ZnO nanowires were grown by molecular beam epitaxy on c-plane sapphire using a procedure reported previously30 (see the SI for details). A TEM image of a typical nanowire (Figure 1a) shows a wurtzite crystal structure with a growth axis along the [0001] direction with atomic planes shown in Figure 1b. The cross section of these nanowires varies from nanowire to nanowire due to local variations on the substrate during growth. Figure 2a shows a high-angle annular dark-field (HAADF) STEM image of a portion of a ZnO nanowire (∼60 nm wide) where a hyperspectral CL image (51 × 10 spatial pixels) was acquired. With uniform composition, the HAADF signal can be taken as a measurement of local thickness, constraining the possible nanowire cross sections. The approximately uniform HAADF intensity across the direction normal to the nanowire shows that its thickness does not vary significantly, consistent, for example, with a near-rectangular section (see Figure SI1). To determine the CL signal generated from defects, the electron beam was held at one spot to locally induce knock-on damage (i.e., vacancies) in the crystal lattice17 for ∼10 s, resulting in a dose over two orders of magnitude higher than CL imaging. This location can be seen in the HAADF image (Figure 2a) as a dark spot approximately half way along the nanowire, as indicated by the arrow. There is a protrusion on the left side of the nanowire that may have been initiated by a defect occurring during growth; in this case, due to the axial growth direction, the defect should be localized toward the bottom of the protrusion, as observed. For other nanowires, we performed HRTEM prior to CL and found that nanowires that had been imaged showed reduced CL intensity in subsequent measurements compared with unexposed structures despite limiting dose and use of a cryo-holder. Conversely, HRTEM performed after STEM-CL acquisition is hindered by amorphous carbon deposition during STEM and potentially other STEM damage artifacts. In less sensitive materials, or with further detector refinement, atomic resolution spatial imaging might be combined with CL; here we focused on maximizing the resolution of the CL maps.

Figure 2b shows three archetypical spectra from the nanowire interior (red), near the base of the protrusion on the left side of the nanowire (green), and just outside the nanowire edge using an aloof beam condition (blue). The interior and protrusion spectra both have a near-band-edge (NBE) emission peak centered at 3.4 eV (370 nm), consistent with the range of band gaps widely reported for both bulk ZnO and ZnO nanowires.35–37 This band gap is in close agreement with the room-temperature PL data (Figure 2m), where increases in PL peak energy with decreasing temperature have been previously reported.38 At lower energies there are small differences between the spectrum collected at the protrusion that has a broad low intensity peak in the range 1.8 to 2.5 eV compared with the bulk spectrum collected far away from any defects visible in the HAADF, for which this peak is either weaker or not present at all. Given the similar intensity in the two spectra from 2.5 to 3.0 eV, this difference is likely to be significant, as confirmed by the further analysis below. The aloof beam spectrum (referred to as S luminescence) is likely caused by surface states and is strongest in the blue, with a peak in counts at −3 eV. The aloof spectrum was fitted with a Gaussian after applying a Jacobian transformation (see Figures SI2 and SI3 for fits), giving a central energy of 2.5 eV, consistent with energies previously reported for surface states.17–21 This range of spectral features (in particular, the differing surface and bulk luminescence) was observed in multiple nanowires, including nanowires from a different growth run and nanowires measured on different days, as shown in Figure SI4.

To investigate the spatial dependence of the different luminescence signals, the CL intensity maps were summed
over consecutive wavelength/energy ranges (Figure 2c−l). By comparing these maps with the HAADF STEM image (Figure 2a), it is clear that the CL intensity varies across the nanowire in a strongly energy-dependent fashion, which must relate to different physical emission processes. On the basis of these maps, there appear to be four distinct spectral components. The UV/purple emission is strongest along the central axis of the nanowire, the blue emission is strongest at the nanowire surface, the green/yellow emission is strongest at the bottom of the protrusion (the likely location of a defect) in the nanowire, and low-energy red emission is present throughout the nanowire. Ensemble PL measures emission due to recombination via the same mechanisms as CL spectra and can therefore be used in tandem to assist interpretation.27

PL measurements on an ensemble of ZnO nanowires grown under similar conditions (Figure 2m) provide high signal-to-noise ratio (SNR) averaged data that confirm the identification of the separate components. The relative amplitudes of the NBE and defect emission peaks differ between the PL (Figure 2m) and CL (Figure 2b). The altered ratio may simply be due to the ensemble averaging in PL measurements of different quality nanostructures (with slightly different luminescence properties) but may also be fundamentally affected by the local excitation rate. The intensity of defect PL (relative to NBE PL) has been shown to decrease with increasing PL pump intensity as defects saturate.39 As such, the high pump intensity of the very narrow STEM beam may cause the reduced amplitude of defect peaks in CL spectra. A study of STEM-CL at reduced beam currents would be interesting but is currently limited by CL detector efficiency. Recent work on SnO2 nanowires23 has also identified differences between PL and STEM-CL spectra; these effects were attributed either to low-quality polycrystalline SnO2 byproducts or more directly to the dependence of the recombination route on excitation mechanisms. In the cited work on SnO2, a radial variation of CL signals (i.e., a reduction in the peak CL intensity at nanowire surfaces) was also reported, analogous to the reduction in ZnO NBE emission reported here. Spectral shifts on length scales of

Figure 2. (a) HAADF STEM image of a ZnO nanowire. A dark spot where the electron beam was held to induce damage is indicated by the red arrow. (b) CL spectra acquired with the electron beam positioned in the nanowire’s interior (red, where the solid line is the dashed scaled by a factor of 15), outside the nanowire (blue), and near the protrusion (green). Line colors correspond to the locations shown by the boxes in panel a. A five-point moving average is applied. Each spectrum is normalized separately for display purposes. (c−l) CL intensity maps generated from a hyperspectral CL image acquired over panel a with a pixel size of 10.2 nm. The brightness in each pixel is proportional to the CL integrated intensity across the wavelength/energy range indicated below each map, as shown in the color bar in panel b. Each map is normalized to the most intense pixel and colored according to the visible spectrum and its respective mean wavelength (UV is shown grayscale). (m) PL spectrum from a ZnO nanowire ensemble. The thick red line indicates a break between two spectra collected using different filters.
10 nm were studied via linear scans across the diameter rather than mapped over the entire structure.

The PL spectrum (Figure 2m) has three obvious peaks: an intense NBE peak, a small peak at energies below the NBE peak (similar to S in CL spectra), and an intense peak in the green significantly more intense than the peak at similar energies seen in CL. The peak in the green requires a pair of Gaussians for a good fit (see Figure SI3), indicating that this lower energy peak is composed of at least two overlapping components and is fit here by two: defect emission 1 and 2 (DE1 and DE2). The central energy of surface luminescence S is close to that previously reported, with a similar breadth.\textsuperscript{17–21} DE1 occurs at 2.1 eV, which has previously been associated with oxygen vacancies.\textsuperscript{14} Emission at 1.8 eV (DE2) has been associated with Zn interstitials.\textsuperscript{40}

Because the emission peaks are broad, each intensity map in Figure 2 includes contributions from multiple overlapping components. Therefore, to understand the correlation between CL emission and nanowire features, the hyperspectral CL image was deconvolved using four spectral components. Each CL spectrum was fitted to a linear superposition of the NBE, S, DE1, and DE2 peaks, where the energy and width of the distributions were fixed according to fits from the CL (NBE and S) and PL (DE1 and DE2) spectra\textsuperscript{41} (see Figures SI2 and SI3 and Table 1). Although room-temperature PL is used to fit the exciton diffusion length, made by polychromatic SEM CL, were \~200 nm in nanowires of 200 nm radius at 5 K\textsuperscript{44} and \~100 nm in a \~50 nm diameter nanowire at 300 K. The NBE component is the dominant contribution to total CL intensity, exceeding the contributions of S, DE1, and DE2. Given the nanowire’s uniform thickness profile (shown in Figure SI1), the surface suppression cannot be a thickness effect and must be related to competing recombination mechanisms. Similarly, alternative recombination routes explain the NBE suppression at the defects. The suppression of NBE emission shows that the exciton lifetime is longer than the lifetime associated with recombination across these defects.

![Figure 3](https://example.com/figure3)

**Figure 3.** (a) HAADF STEM image of nanowire repeated from Figure 2a for reference. Scale bar is 50 nm. Amplitude maps of (b) near-band-edge emission, (c) surface emission, (d) defect emission 1, and (e) defect emission 2 deconvolved components. (f) Counts integrated across all energies. The intensity in each map is normalized and smoothed for presentation clarity by applying a Gaussian blur of pixels with a radius equal to half the CL pixel size (5.1 nm). To generate maps b–e, CL spectra are transformed to counts/eV and then fit. The amplitudes of these maps are 42 560, 2837, 2533, and 2981 respectively.

**Table 1.** Fit Results for Central Energy ($E_0$) and Peak Width ($\sigma$) of the Four Components Used To Deconvolve CL Spectra\textsuperscript{a}

| Component | PL/CL   | $E_0$ (eV) | $\sigma$ (eV) |
|-----------|---------|------------|---------------|
| NBE       | CL (103 K) | 3.40 ± 0.01 | 0.106 ± 0.001 |
| S         | CL (103 K) | 2.53 ± 0.05 | 0.61 ± 0.05   |
| DE1       | PL (300 K) | 2.08 ± 0.01 | 0.194 ± 0.001 |
| DE2       | PL (300 K) | 1.82 ± 0.01 | 0.091 ± 0.001 |

\textsuperscript{a}Second column denotes whether CL or PL data were used to fit the peaks and the temperature at which these measurements were performed. Fits were performed on data transformed into counts/eV (see the SI for details).
interacting with the nanowire) is likely due to the excitation of surface plasmons,\(^{18,45}\) which decay to excitons and subsequently recombine via surface traps in the ZnO, but may also be caused by stray electrons that directly excite excitons preferentially at the nanowire surface. The peak of S intensity extends into the nanowire due to excitons excited in the nanowire diffusing to the surface, where they subsequently decay. The yellow/green defect emission, DE1 (Figure 3d), is strongest near the two structural defects in the nanowire, the same regions where NBE is less intense. Far away from structural defects, the DE1 emission is nonzero, indicating that this trap is present throughout the nanowire but at higher concentrations around structural defects. DE1 is centered at 2.1 eV, and luminescence at this energy has previously been attributed to O vacancies,\(^{14}\) consistent with vacancies induced by knock-on damage. DE1 is less intense at the surfaces of the nanowire, indicating that DE1 is slower than recombination due to surface states. The orange/red defect emission, DE2 (Figure 3e), has a central energy of 1.8 eV and is approximately uniform along the nanowire, with a small decrease at the nanowire surface, and occurs at energies previously associated with Zn interstitials,\(^{40}\) although others\(^{40}\) question whether Zn interstitials are stable in ZnO.

The total integral of the CL counts, mapped across the nanowire (Figure 3f), highlights the occurrence of nonradiative recombination. In a nanowire with approximately uniform thickness, the naive assumption would be that counts should be generated uniformly across the nanowire. Instead, it is clear that the integrated counts drop both at the edge of the nanowire and in the vicinity of peaks in DE1. This trend indicates that both the nanowire surface and defects within the nanowire provide nonradiative recombination routes, in addition to the radiative recombination generating CL counts associated with these features. Quantum efficiency drops at regions of increased visible luminescence (defects), a poor prospect for optical ZnO devices operating in the visible spectrum, as nonradiative processes will reduce photoemission.

High-resolution STEM-CL mapping of ZnO nanowires can resolve variations both axial and perpendicular to growth axis, allowing structural features and luminescence signals to be related. This powerful characterization technique provides nanoscopic insight into luminescence processes and how they correlate locally to defects. In this Letter, we present high-resolution hyperspectral maps of a single nanowire by STEM-CL. However, simple mapping of the CL emission at different energies does not distinguish all of the underlying processes because their luminescence signatures overlap. Ensemble PL measurements can help to identify the key spectral features present within the sample (i.e., peak energy and spectral width), allowing them to be spectrally deconvolved from the noisier CL data. Maps of these components across a single nanowire identify an exciton diffusion length of \(\sim 50 \text{ nm}\) and show that the dominant NBE luminescence is confined to the core of the nanowire and is suppressed near the surface (S) or defect sites (DE1, most likely zinc interstitials). Because the integrated intensity drops both at the surface and close to regions where DE1 is particularly bright, nonradiative recombination must occur at these locations at increased rates. A hierarchy of time scales (including trapping and recombination processes) can be established by considering the local suppression of different luminescence mechanisms. The exciton recombination timescale is longer than time scales associated with DE1 emission, which are, in turn, longer than time scales associated with S luminescence. These different time scales arise from different lifetimes and trapping times associated with different luminescence processes and can be measured by complementary techniques such as transient absorption spectroscopy.

Spatial information about the distribution of defects will aid detailed simulations of ZnO nanostructures, providing a better understanding of consequent device properties. For example, band structure calculations may shed light on transport or photocatalysis, whereas the spatial distribution of optically active defects may inform how such nanostructures are deployed as optical gain media. The general STEM-CL approach may be more widely applied in the future to study other nanowire systems developed for optoelectronics or photocatalytic applications, in particular, those with in-built heterostructures or junctions. Improvements in optical detection will increase the spatial resolution of the CL signal and reduce exposure times. Future measurements may combine atomic-resolution TEM or STEM imaging of defect structures or other analytical techniques with high-resolution CL to establish definitive assignments that, despite significant work, are still controversial in ZnO.

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