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Simultaneous two-color lasing in a single CdSSe heterostructure nanosheet

F Fan, Z Liu, L Yin, P L Nichols, H Ning, S Turkdogan and C Z Ning

School of Electrical, Computer, and Energy Engineering, Arizona State University, Tempe, AZ 85287, USA
E-mail: cning@asu.edu

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Abstract
The ability of a single monolithic semiconductor structure to emit or lase in a broad spectrum range is of great importance for many applications such as solid-state lighting and multi-spectrum detection. But spectral range of a laser or light-emitting diode made of a given semiconductor is typically limited by its emission or gain bandwidth. Due to lattice mismatch, it is typically difficult to grow thin film or bulk materials with very different bandgaps in a monolithic fashion. But nanomaterials such as nanowires, nanobelts, nanosheets provide a unique opportunity. Here we report our experimental results demonstrating simultaneous lasing in two visible colors at 526 and 623 nm from a single CdSSe heterostructure nanosheet at room temperature. The 97 nm wavelength separation of the two colors is significantly larger than the gain bandwidth of a typical single II–VI semiconductor material. Such lasing and light emission in a wide spectrum range from a single monolithic structure will have important applications mentioned above.

1. Introduction

Realization of monolithic semiconductor structures with multiple or a wide range of bandgaps have many important applications in solid-state lighting, high efficient solar cells and multi-spectral detection. In solid-state lighting applications, a single semiconductor structure or device capable of emitting or lasing in the entire visible range (400–700 nm) would have a fundamental impact on the ultimate solution of efficient lighting. It was recently demonstrated [1] that four lasers of discrete spectral lines are virtually indistinguishable from state of the art continuous-spectrum white reference illuminant, thus demonstrating the feasibility of laser lighting. In addition, lasers can be much more efficient than light-emitting diodes (LEDs). Similarly, discrete multicolor lasers have many other applications such as full-color laser display [2], white laser illumination [1] and biological inspection [3]. Even though the experimental demonstration was carried out with four separate lasers [1], a much more preferred approach would be to have all lasers of different colors produced on a same platform in a compact fashion. But achieving such emission or lasing in a wide spectrum is fundamentally challenging for several reasons. First, typical semiconductors of a given composition under reasonable excitation conditions have a gain bandwidth less than 30 nm [4–12]. Wavelength tunability of a single mode, or mode separation of multiple modes, is thus limited by such a bandwidth. Second, to have emission bandwidth significantly broader than that of a single-composition material, heterogeneous integration or growth of different semiconductors is necessary. Due to the large variation or significant mismatch of the lattice constants for semiconductors with distinct material compositions, growth of such structures on a single substrate becomes extremely difficult. Finally, even such materials could be grown, but it is still a challenge to design an appropriate cavity structure to support wavelengths that are widely separated. This is because the short-wavelength light will be strongly absorbed
by the narrow gap semiconductors involved, such that only the long-wavelength light can eventually reach the threshold and become lasing. Various attempts have been made recently to overcome these challenges and to extend the spectral range of light emission and lasing beyond the gain bandwidth of a single material. Multi-color or dual-color lasing has been observed by placing multiple nanowires within a microfiber [13], in multi-section quantum dot distributed feedback lasers [14], or by using fast-switching of different dyes in a microfluidic channel [15]. However these multi-color lasers are either achieved in separated devices, or with wavelength separation still limited within the gain bandwidth of a single semiconductor material.

In this paper, we report the first experimental observation of simultaneous lasing in two visible colors (at 526 and 623 nm, respectively) from a single monolithic semiconductor structure. The wavelength separation of two lasing colors of 97 nm is much larger than the gain bandwidth of a single composition semiconductor material. The unique structure in this paper that enabled such dual wavelength lasing is a heterostructure nanosheet or nanobelt of the ternary semiconductor CdSSe produced in a single monolithic growth. Our results were enabled by some unique quality of nanowires and nanobelts that have been the focus of light emission and lasing research for the last 10 years [16–24]. Semiconductor nanowires and nanobelts grown using the metal-catalyzed vapor–liquid–solid mechanism have opened new possibilities for multi-color emission or lasing [17–22]. Even though semiconductor nanoparticles can also achieve multi-color emissions, but it is typically challenging to spatially address or control each individual color, since they are typically mixed uniformly. In addition, eventual electrical injection is needed for such applications and the poor conductivity in nanoparticles make electrical injection very inefficient. One of the important features of nanowires and nanobelts for our purpose is the ability to grow different materials with different lattice constants without introducing too many defects to degrade optical properties. This property has been utilized in the past to grow materials with very different or graded alloy compositions [18–21] to achieve light emission in a wide range, mostly from different wires and more recently from a single wire [25]. While we recently have demonstrated multi-color lasing from many wires on a single nanowire or belt has not been demonstrated, especially when the wavelength separation is wider than the gain bandwidth.

2. Sample growth and characterization

The CdSSe lateral heterostructure nanosheets were grown through a chemical vapor deposition (CVD) approach using CdS and CdSe powders (Alfa Aesar, 99.99% purity) as source materials. Si substrate with 10 nm Au film as catalyst was used for the growth. The reactor was first evacuated to a pressure below 0.2Torr by a mechanical pump. 150 sccm N2 was then introduced into the system, and flow was maintained for 30 min prior to growth. In the first step of the growth, only CdS powder was thermally evaporated at 880 °C for 1 h. Subsequently, CdSe source was introduced into the furnace, while the temperature was gradually lowered to 840 °C and held for 40 min to grow the CdSe-rich sections. The growth processes is similar to that of [26], but we reversed the growth sequence to grown CdS first. Such sequential growth ensures a wide CdS-rich center section surrounded by CdSe-rich edges (see figure 1(B)). The large width of CdS-rich sections is especially significant as it needs to be wide enough to achieve sufficient optical confinement for the shorter wavelength lasing modes.

The composition distribution on the CdSSe heterostructure nanosheet is shown in figure 1. Figure 1(A) shows the real color photoluminescence (PL) image of the heterostructure nanosheet under 405 nm continuous-wave (CW) laser excitation. This nanosheet has width of 11.2 μm and length of 41.4 μm. As seen in the image, the nanosheet consists of three stripes with green emission in the center (grown initially with CdS-rich composition) and red/orange emission on both edges (grown subsequently with CdSe-rich composition). With proper end facets, the three stripes can form three coupled cavities capable of supporting various propagating modes along the length direction. Of particular interest are the two wider stripes (the green one in the center and the red one on the right side) which form two rectangular, coupled cavities in a side-by-side arrangement, as also schematically shown in figure 2(C). We performed micro-PL measurements by scanning the focused laser beam with a 1 μm across the width of the nanosheet along the dashed line in figure 1(A). The PL peak positions vary from green emission at 559 nm to orange emission at 602 nm with a transition occurring roughly at 3 μm in figure 1(C). Assuming the PL emission is from ternary CdSxSex1−x, we use the interpolation formula [27]

\[ E_g (\text{CdS}_x \text{Se}_{1-x}) = xE_g (\text{CdS}) + (1 - x)E_g (\text{CdSe}) \]  

Figure 1. (A) Real-color PL image of a CdSSe lateral heterostructure nanosheet excited by 405 continuous-wave laser. (B) TEM image of the same nanosheet. The scale bars in (A) and (B) are 10 μm. (C) Comparison of sulfur molar fraction between EDS scan (hollow triangles) and micro-PL scan (solid squares) along the white dashed line in (A) and (B).
to fit the PL peak and to extract the alloy composition. The result is plotted in figure 1(C), where the sulfur fraction along the scanning path with transition from 45% to 67%.

To further confirm the composition distribution across the heterojunction, the energy dispersive x-ray spectroscopy (EDS) in a transmission electron microscope (TEM) is also performed along the same path as the PL scan. Figure 1(B) shows the TEM image of this nanosheet. The image was collected with a JEOL JEM-2010 High-Resolution TEM at 200 kV, equipped with a Link EDS detector. The sulfur compositions obtained from EDS line-scan along the dashed line are plotted in figure 1(C), consistent with the values from micro-PL scanning.

3. Experimental and simulation results

Figure 2(A) shows a PL image of another lateral heterostructure nanosheet. The dimensions of this nanosheet are 25.0 μm (length) by 17.4 μm (width) by 200 nm (thickness). The nanosheet was cleaved from a longer piece by the bend-to-fracture method [28]. With this procedure, high-quality end facets were created to serve as facet reflectors and to define the laser cavities. The widths of the CdS-rich (green) and CdSe-rich (orange) sections are 14.8 and 2.6 μm, with PL emission peaks at 546 (green) and 591 nm (orange), respectively. As shown in the PL image, there are broken nanosheet debris surrounding the nanosheet of interest, observable by the lighter orange areas within the nanosheet and the excess green areas outside of the nanosheet perimeter. To demonstrate the capability of dual-color lasing, the sample was transferred onto a sapphire substrate and loaded into a cryostat for low temperature measurement. We optically pumped the nanosheet by the third harmonic of a Q-switched Nd:YAG laser (355 nm, 10 Hz, 9 ns). The laser was focused into a 120 μm by 60 μm elliptical beam at an angle of 60° from the sample normal direction. The real-color image of single pulse induced lasing at higher pumping level at 77 K is shown in figure 2(B). As can be seen clearly, the largely uniform spontaneous emission of figure 2(A) is now replaced by the green and orange bright spots along the end facets, signifying the occurrence of lasing. Due to the gain-guiding effect and the refractive index contrast, green and orange emission can be well-separated and confined in their own cavities, which is schematically shown in figure 2(C).

This is reflected in figure 2(B), where green and red emissions from the end facets are from the respective segments of CdS-rich or CdSe-rich materials. The interaction between the green and orange modes is minimized in this structure. The side-by-side arrangement is an important geometry for reaching the respective lasing thresholds of both colors. This is in contrast to structures where bandgap changes along the length of a nanowire [29], where short-wavelength emission from the wide bandgap region (CdSe-rich) can be significantly absorbed by the narrow bandgap region (CdS-rich). Such absorption will increase the threshold for the short wavelength lasing, very often to a degree that is too high to achieve lasing. Our side-by-side arrangement minimizes such undesired absorption and allows the respective lasing thresholds of both wavelengths to be reached.

Figure 2(D) shows the PL spectra under excitation of a single pulse with increasing pump power density. At lower pumping levels, only broadband spontaneous emission from the CdS-rich and CdSe-rich sections is observed. With increasing pump pulse power, a narrow peak appears at 592 nm at an excitation level of 173 kW cm⁻², indicating the onset of lasing for the orange color in the CdSe-rich section. The same is observed for green color lasing at 534 nm in the CdS-rich section when the excitation level reaches 241 kW cm⁻². The 58 nm separation of the two lasing peaks is much larger than the gain bandwidth of a typical Zn-Cd-S-Se semiconductor material system including CdS or CdSe [2–8]. The narrowest linewidths of the two colors in figure 2(D) are around 4 nm, which are significantly broader than the typical linewidth of a single mode semiconductor laser. Detailed examination proves such broad linewidths of both sections under higher excitation intensities is due to the well-known multimode lasing behavior [30–35] and spectrometer resolution limitation (at 0.1 nm). To confirm this, we performed high resolution measurement for the orange lasing peak and compared with the low resolution measurement (shown in figure 3) to examine the detailed mode features. The high resolution measurement reveals that the wide peak in low resolution measurement actually consists of multiple peaks with individual linewidths as narrow as 0.1–0.3 nm. The multiple lasing peaks can be divided into four groups and each group consists of a series of closely spaced peaks. The four groups of lasing peaks can be identified as the longitudinal modes of different orders with each group consisting of several closely spaced transverse modes. The wavelengths of three transverse modes TE₉₀, TE₉₀, TE₃₀ and the corresponding longitudinal modes are also calculated and labeled in figure 3 for comparison. The mode spacing between
the transverse modes is typically smaller than the longitudinal mode separation and can be as small as 0.2 nm, which almost reaches the resolution limitation of our spectrometer system. We attribute the multimode lasing features to the large size of the nanosheet structures in both directions, as compared to the wavelengths. With increasing pump intensity, more modes are excited above the threshold (see appendix for another nanowire case). Their close proximity to each other results in broader excited above the threshold (see appendix for another nanowire case). Their close proximity to each other results in broader

In order to demonstrate spatial separation of the green and orange lasing modes in the side-by-side cavities of the heterostructure nanosheet, we performed a 2D simulation for the optical modes at the lasing wavelengths of 534 and 592 nm, as observed experimentally (see figure 2). In this 2D model, the length of the nanosheet is assumed to be infinitely long with the propogation direction perpendicular to the paper plane. The figure shows the cross section of a nanosheet in the thickness-width plane. The propagation direction is perpendicular to the paper plane. (A) The schematic of the nanosheet cross-section. The green (left) and red (right) rectangles represent the CdS-rich (14.8 μm width) and CdSe rich segments (2.6 μm width), respectively. The thickness is 200 nm. (B–D) The first three modes around 534 nm. (E–G) The first three modes around 592 nm.

The green, red and blue vertical lines represent the wavelengths of their PL emission peaks are (cross-section as shown in figure 2). The imaginary part of the complex index of refraction, \( \lambda \) is the wavelength and \( g \) is the gain or absorption coefficient (with a negative sign). A positive extinction coefficient represents gain, and negative represents absorption. The absorption of CdS0.49Se0.51 at 534 nm is 50 000 cm\(^{-1}\) based on the calculations and data in [38]. Under optical pumping, the gain coefficients in active regions are assumed to be 3000 cm\(^{-1}\) which is a reasonable assumption to overcome the mirror loss.

From the simulation, the first three modes with highest gain are selected for each wavelength. The amplitude of the electric fields of these modes at 534 and 592 nm are shown in figure 4(B–D) and (E–G), respectively. For both wavelengths, the highest-gain modes are well-confined in their respective active regions (CdS-rich segment for 534 nm, on the left in figure 4(A), and CdSe-rich segment for 592 nm, on the right in figure 4(A)). It is important to determine the confinement factor, defined as the ration of the modal gain to the material gain [39, 40], as it is known now that it can be significantly large in nanowires and nanobelts [30, 39, 40]. The confinement factors for all of these modes are slightly larger than unit at 1.03. In general, modes of both wavelengths can be confined in either the CdS-rich or CdSe-rich segment due to the contrast between the refractive indices. However, only the modes in appropriate gain medium with wavelength inside the gain bandwidth have the possibility to lase. As a result, high gain modes at 534 nm (or 592 nm) can only lase in the CdS-rich (or CdSe-rich) active regions. The refractive index contrast and the gain confinement ensure that the modes of both wavelengths are confined within their respective gain media without much leakage to the lossy regions. This is especially important for the green modes, so that they will not be subject to too strong absorption by the narrow gap material of the CdSe-rich side. The simulation proves our previous statement that the CdSSe heterostructure nanosheet provides an appropriate structure to support dual-wavelength lasing in a single structure.
The integrated light intensity is plotted as a function of pumping power density on both log–log (figure 5(A)) and linear scales (figure 5(B)) to corroborate the spectral features and to further establish the lasing behavior at two distinct wavelengths. Shown in figure 5(B), the lasing intensity of both colors increases linearly with pump power after threshold (260 kW cm\(^{-2}\) for the green modes and 230 kW cm\(^{-2}\) for the orange modes). The total spontaneous emission output increases much more slowly than the lasing modes in the high excitation region, demonstrating the typical behavior associated with density clamping (or slowly increasing) near the lasing threshold. The log–log scale of the lasing output as a function of the excitation power features the well-known S-like behavior with three significantly different regions with different slopes, as indicated by various straight lines used to extract the slopes. At low pumping, the spontaneous emission is dominant, so the mode intensities increase linearly with a slope of 1.1 for orange and 1.3 for green. Around the thresholds, the output intensities of both lasing modes show superlinear increases with slopes of 5.1 for orange and 4.7 for green. With further increase of the excitation power, the slopes of both curves return to 1.1, indicating the above-threshold lasing operation of both colors. The dependence of total intensity on pumping displays a typical threshold behavior of multimode lasing, which is somewhat more complicated than that of a single mode laser. To examine the lasing threshold behavior including the multiple transverse and longitudinal modes, experimental data of total light intensity as a function of the excitation power features the well-known S-like behavior including the multiple transverse and longitudinal modes. These transverse modes contribute to the spontaneous emission factor \(\beta\) of each lasing mode. For a semiconductor nanowire laser, \(\beta\) is less than 0.001 [35]. For a semiconductor nanowire laser, \(\beta\) becomes larger, around 0.02 [30, 33]. Figure 5(C) shows the fitting result. The best fitting was obtained when \(x_0\) was taken as 0.085 for green mode and 0.090 for orange mode. The relatively large \(\beta\) as compared to nanowire lasers can be understood by the relationship between \(x_0\) and spontaneous emission factor \(\beta\). \(x_0 = 0.02\) was obtained when a clear set of longitudinal lasing modes was observed from the nanowire. In our case, the multimode lasing from the nanosheet contains transverse in addition to longitudinal modes. These transverse modes survive during the lasing process due to the large confinement factor, as previously addressed. All of these modes contribute to the spontaneous emission factor \(x_0\) to increase its magnitude. As a result, the light-in-light-out curve of nanosheet laser shows a slightly soft threshold.

Room temperature (RT) lasing has also been achieved on a larger nanosheet (66.5 \(\mu\)m in length, 33.0 \(\mu\)m in width and 200 nm in thickness). As shown in figure 6(A), the widths of the green and the wider orange stripes are around 30 and 3 \(\mu\)m, respectively. The real color lasing image and PL spectra evolution are shown in figures 6(B) and (C), respectively. The lasing wavelengths of the orange and green peaks are 613 and 562 nm, respectively, with a wavelength separation of 51 nm. The estimated threshold is 300 kW cm\(^{-2}\) for the green modes and 227 kW cm\(^{-2}\) for the orange modes. The lasing sequence for the two colors at RT is consistent with the result at 77 K, both exhibiting higher thresholds for the green modes than for the orange modes. As in figure 2, the main peaks shown in figure 6(B) broaden as pumping power is increased due to the excitation of increased number of modes, as we discussed in connection with figure 3.

The wavelength separation of two lasing colors can be further expanded by improving the growth methods of CdSSe nanosheet. The improved growing steps are the same as...
described in section 2 except that we totally terminated the CdS source supply before introducing CdSe source. By doing this, the ions exchange between sulfur and selenium can be minimized during the second step of the growth and as a result, the wavelength separation of two color lasing can be further expanded. Figure 7 demonstrates the two color lasing results of the nanosheet by changing the growth method. The lasing wavelengths are 526 and 623 nm, respectively with a separation of 97 nm. The longer wavelength has now been fully expanded into the red color range. The dimensions of the nanosheet in figure 7(A) are 23.5 μm (length) by 5.2 μm (width) by 200 nm (thickness), much smaller than that of the nanosheets demonstrated above. As a result, less significant multimode behavior is shown in figure 7(C). Other features are similar to the previous ones.

4. Conclusion

To summarize, we demonstrated simultaneous lasing in two distinct visible colors from a single CdSSe lateral heterostructure nanosheet. This demonstration was made possible by utilizing an important progress made recently in the growth of nano-materials using the metal-catalyzed vapor–liquid–solid CVD growth. The key material enabler is the virtually substrate-insensitive growth of heterostructures with wide composition variation within a single nanosheet. The side-by-side geometry overcomes one of the most challenging issues with multi-color lasing: the absorption of short-wavelength light by the narrow gap semiconductors. The dual-color lasing was established by a combination of different measurements including high and low resolution spectral measurements, integrated light intensity measurements at various pumping levels, and changes of near-field spatial profiles with pumping levels. The multimode lasing behavior has been clearly shown in high resolution measurements. In low resolution measurements, the observed linewidth broadening with increasing pump power is attributed to the increasing number of modes contributing to the overall peak width. This has been known for multimode lasing in semiconductors [30–35, 40]. One of the key quantities is the wavelength separation between the two colors, which is eventually expanded to 97 nm at RT, much larger than the typical gain bandwidth of semiconductor materials (<30 nm) in this wavelength range. Lasing with such widely separated colors from a single semiconductor structure can be important for many applications.

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Appendix. Experimental illustration of modes evolution with pumping

To understand the increasing linewidth with pumping as measured with low resolution spectroscopy, we studied a CdS nanowire with multiple transverse and longitudinal modes using a high spectral resolution measurement, with the results presented in figure A1. Since our nanowire has a much shorter length (at 10 μm) than the lengths of those nanosheet structures studied earlier, the longitudinal modes are separated...
with larger spectral intervals. Therefore, it is much easier to distinguish each longitudinal mode. Similar to the lasing spectrum of the nanosheet shown in figure 3, the separated groups of modes represent different longitudinal modes and the fine peaks in each group are different transverse modes. When raising the pumping power, more longitudinal modes appear at longer wavelength side, from 1 in the bottom panel to 5 in the top panel. This can be explained by the gain spectrum broadening and heating effect. The linewidth of each longitudinal mode increases with pumping owing to the increasing numbers of transverse modes. This multimode lasing and linewidth broadening effect was observed in both nanowire and nanosheet [30, 41, 42].

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