Photopumped laser oscillation and charge-injected luminescence from organic semiconductor single crystals of a thiophene/phenylene co-oligomer

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We have demonstrated that single crystals of a thiophene/phenylene co-oligomer [α, ω-bis-biphenyl-4-yl-terthiophene (BP3T)] show interesting photonic aspects: (1) the self-waveguided amplified spontaneous light emissions with a comparable low threshold of 8 μJ/cm² to other optimized organic solid-state laser systems, and (2) the laser oscillation based on the optical self-confinement effect in the crystals. We have also presented electroluminescence from the crystals based on bipolar injection and the crystals’ tolerance for intense current driving. These achievements strongly imply that BP3T crystals are a promising candidate for organic laser diodes. © 2005 American Institute of Physics. [DOI: 10.1063/1.2138361]

The achievement of long-lasting, bright electroluminescence with high efficiency in both polymeric and low-molecular-weight organic materials has inspired the development of a new class of laser diodes (LDs). Within this development, optically induced solid-state organic lasers have been proposed and fabricated based on thin films by methods such as solution casting and vacuum deposition with a wide range of related organic semiconductors. As a counterpart of these research streams, crystalline molecular materials also get much attention as promising candidates for organic LDs demanding high-density electric pumping for population inversion. Organic molecular single crystals exhibit superior charge-carrier transport properties to amorphous and polymeric organic semiconductors. This advantage is due to the inherent long-range structural ordering of these single crystals. Furthermore, there are several examples of stimulated emission, including amplified spontaneous emission (ASE), from molecular single crystals. These excellent photonic and electronic features imply the possibility of current-injection lasing. However, charge-injected luminescence from organic single crystals with high stimulated emission properties has not yet been reported. We have achieved optically induced laser oscillation and charge-injected luminescence from a molecular single crystal. This breakthrough opens up a new research path toward developing organic LDs.

The synthesis of a thiophene/phenylene co-oligomer, α, ω-bis-biphenyl-terthiophene (BP3T) used can be seen in our previous work. The procedures of the crystal growth were also described in detail elsewhere. The charged pressure of the N₂+H₂ (10 mol %) growth gas was −18.6 kPa (a differential pressure from the atmosphere) at room temperature. The temperature of the growth cylinder was maintained at 360 °C at the material-loaded position (bottom) and 300 °C at the growth position (top) during crystal growth (about 150 h). The melting temperature of BP3T is 388 °C. The source materials of the crystals were highly purified by sublimation in advance. A N₂ gas laser that generated a light pulse of 500 ps duration at 337 nm was used as an excitation light source for the gain-narrowing experiment. The detailed experimental setup was described in our previous work.

Fabrication and measurement of the charge-injected luminescence device were carried out follows. A gold electrode that worked as an anode was thermally deposited in the thickness of 15 nm through a shadow mask on a polished glass substrate. Referring to our previous work, the thickness of the gold layer was determined to balance electric conductivity and adhesivity of crystals to both the glass substrate and gold electrode. A selected single crystal, of thickness below 1 μm, was placed on the gold electrode, and then the crystal spontaneously adhered to the electrode and substrate. Finally, another electrode as a cathode of Au, Al, or Li was thermally deposited on the fixed crystal through a shadow mask. For Li, the thickness of the metal layer was 10 nm, and then a 200-nm-thick Al layer was simultaneously deposited on the Li layer for passivating the active Li under ambient conditions. For Au and Al, a 200-nm-thick layer of the metals was used as the upper electrode. The current-voltage characteristics of the devices were measured under a vacuum (10⁻³ Pa or less) at room temperature. The electroluminescence (EL) spectra were measured with a spectrometer (Ocean Optics, USB2000).

We took an optical photograph (Fig. 1(a)) and a scanning electron microscope image (Fig. 1(b)) of BP3T, thin-plate-like crystals prepared with the vapor phase growth method using buffer gas. The figure also shows the chemical structure of BP3T (the inset of Fig. 1(a)). As the optical photograph shows, the BP3T crystal is a smooth-surfaced single platelet about 1 × 1 mm². Fluorescence was observed only at the crystal edges except for a couple of attached small crystals. The micrograph observations of the crystals under crossed Nicols configuration indicate that bright- and dark-field images are each switched at 45° rotation, indicating that the grown crystals are formed with single domain only. That is, the crystals are single crystalline. The electron microscope image (Fig. 1(b)) shows that the surface of the crystal is smooth and flat. The uniform flatness of the crystals provides optical confinements and slab-mode propagation in crystal
and highly electronic contacts with physical adhesions for fabrication of electricity-driven devices.

Figure 2(a) shows the photoluminescence (PL) spectra of BP3T thin-plate-like crystals under weak and intense excitation. The PL spectra narrowed under intense excitation. As plotted in Fig. 3, PL peak intensity increased nonlinearly above excitation laser intensity of 8 μJ/cm², and the spectral width [full width at half maximum (FWHM)] abruptly decreased around the same excitation intensity. These spectral behaviors confirm that the narrowed PL spectra results from gain narrowing, and the threshold for stimulated emission is only 8 μJ/cm². This threshold is comparable to that of other optimized organic laser systems, such as DCM2:Alq and DSB:TPD. The photoemission was highly directional and exhibited superior output stability after thousands of shots with the excitation pulse laser. The ASE wavelength of 618 nm agrees with that of the second vibronic peak (0-1 transition) of the PL spectrum of the BP3T crystal under weak excitation, since the weak shoulder observed at around 580 nm corresponds to the first vibronic peak of the luminescence. Spontaneous emissions (i.e., PL in the weak excitation regime) from conjugated materials are usually accompanied by vibronic peaks where the emission intensities are maximal, producing the largest gain. The relevant vibronic levels are thus sufficient for population inversion in stimulated emission. Thus, BP3T crystals are one of the promising candidates for organic solid-state laser and organic laser diodes. The nonsaturated output of their PL peak intensity until the excitation fluence of 400 μJ/cm² is a particularly useful characteristic.

Figure 2(b) shows the laser oscillation photoluminescence spectrum with high wavelength resolution under intense excitation from another BP3T crystal that has a pair of parallel naturally formed facets that worked as a Fabry-Perot optical resonator. The spectrum has regularly ordered progressive peaks whose interval and FWHM are 104 and 38 pm, respectively. In addition, the center wavelength of the progressive peaks of about 618 nm agrees with the ASE wavelength. These features strongly indicate that the self-cavity laser oscillation occurs in multimode, a result that agrees well with our recent report on optical pumped self-cavity laser oscillation in multimode, a result that agrees well with our recent report on optical pumped self-cavity lasing on crystals of another oligothiophene derivative. Thus, the BP3T crystals confined enough photoluminescent light, and loss caused by absorption and scattering in the crystal and at its surfaces was sufficiently small. Therefore, BP3T crystals work as a self-cavity lasing medium.

Figure 4 shows the current density versus the applied electric field (J-E) characteristics of the charge-injected luminescent devices using BP3T single crystals, of which the device structure was glass substrate/Au (anode)/BP3T crystal/cathode (Au, Al, or Li), and where the electric field is defined as a ratio of the applied voltage to the thickness of the crystal used. As shown in the figure, the Au cathode device shows a reversible J-E characteristic as the reflection of the symmetric device structure. This reversible characteristics strongly indicates that the electric junction between the under-Au electrode and the BP3T crystal, which was fabricated by a simple procedure described before, works as well as a metal deposited junction. Some difference was observed between hole injection characteristics of adhesion electrodes.
Au anode. Inset different metal cathodes. The two characteristics for the Au cathode device are 
transition dipole moment of such molecules is parallel to the
occupied molecular orbital: 2.76 eV, lowest unoccupied molecular orbital: 5.08 eV, 
area may have caused the difference.

FIG. 4. 

<table>
<thead>
<tr>
<th>Electric Field (10^8 V/cm)</th>
<th>Current Density (A/cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>10</td>
<td>1</td>
</tr>
<tr>
<td>20</td>
<td>2</td>
</tr>
<tr>
<td>30</td>
<td>3</td>
</tr>
</tbody>
</table>

slope = 1

long axis, luminescence is emitted around the ab plane. Hence, the crystal radiates most of its EL from the edge face. In addition, since the crystals are thinner than the wavelength of the light, the crystals also worked as waveguides.

In this letter we demonstrated optoelectronic features of a BP3T single crystal. The BP3T crystals exhibited an excellent stimulated emission property; the crystals showed the self-waveguided ASEs with a comparable low threshold of 8 μJ/cm² to other optimized solid-state laser systems, and the crystals with naturally formed parallel facets that worked as an optical resonator showed multimode laser oscillation under optical excitation. Moreover, we also demonstrated EL from the BP3T crystals. The EL was waveguided in the crystal and emitted from the crystal edge face. In addition, BP3T crystals can endure intense electric excitation up to 15 A/cm² at unpulsed dc operation. These optoelectronic features of BP3T crystals clearly reveal that they are a promising candidate for organic LDs.

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8) For instance, a charge carrier mobility as high as ~15 cm²/Vs, exceeding that of a-Si:H and comparable to that of polycrystalline indium-tin-oxide thin films, was reported by V. C. Sundar et al. in molecular crystals of rubrene [V. C. Sundar, J. Zaumseil, V. Podzorov, E. Menard, R. L. Willett, T. Someya, M. E. Gershenson, and J. A. Rogers, Science 303, 1644 (2004)].