Atomically Thin MoS$_2$: A New Direct-Gap Semiconductor

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The electronic properties of ultrathin crystals of molybdenum disulfide consisting of $N = 1, 2, \ldots, 6$ S-Mo-S monolayers have been investigated by optical spectroscopy. Through characterization by absorption, photoluminescence, and photoconductivity spectroscopy, we trace the effect of quantum confinement on the material’s electronic structure. With decreasing thickness, the indirect band gap, which lies below the direct gap in the bulk material, shifts upwards in energy by more than 0.6 eV. This leads to a crossover to a direct-gap material in the limit of the single monolayer. Unlike the bulk material, the MoS$_2$ monolayer emits light strongly. The freestanding monolayer exhibits an increase in luminescence quantum efficiency by more than a factor of 10$^4$ compared with the bulk material.

As a consequence of these different scaling properties and as suggested by previous calculations [8,9], the MoS$_2$ crystals exhibit a crossover from an indirect- to a direct-gap semiconductor in the monolayer limit. In addition to the signatures of this effect in the absorption and photoconductivity spectra, the PL quantum yield (QY) showed a dramatic enhancement in going from the dark, indirect-gap bulk crystal to the bright, direct-gap monolayer [23]. For our suspended samples, we observed an increase of the PL QY by more than a factor of 10$^4$ for the monolayer compared with the bulk crystal.

We prepared mono- and few-layer MoS$_2$ samples from bulk MoS$_2$ (SPI Supplies) by using a mechanical exfoliation technique similar to that employed for graphene [21]. Typical samples ranged from 25 to 200 $\mu$m$^2$ in size. Suspended samples were obtained by depositing the

![Figure 1](https://example.com/figure1.png)

**FIG. 1 (color online).** Lattice structure of MoS$_2$ in both the in- and out-of-plane directions and simplified band structure of bulk MoS$_2$, showing the lowest conduction band $c1$ and the highest split valence bands $v1$ and $v2$. $A$ and $B$ are the direct-gap transitions, and $I$ is the indirect-gap transition. $E_{g1}^i$ is the indirect gap for the bulk, and $E_{g2}^i$ is the direct gap for the monolayer.
MoS$_2$ layers on oxide-covered Si substrates that had been prepared with arrays of holes (1.0 and 1.5 $\mu$m in diameter) [24]. We identified areas with thin MoS$_2$ samples with an optical microscope [Fig. 2(a)]. PL images were also collected to assess sample quality [Fig. 2(b)]. The reflectance contrast of the samples permits determination of their thickness with monolayer accuracy, as confirmed by atomic-force microscopy.

Optical measurements by absorption, PL, and photoconductivity spectroscopy were performed on mono- and few-layer MoS$_2$ samples. All optical measurements were carried out under ambient conditions at room temperature by using an inverted microscope (Nikon) coupled to a grating spectrometer with a CCD camera. The optical beams were focused on the sample with a spot diameter of $\sim$1 $\mu$m. For absorption measurements [25], samples on transparent fused quartz substrates were studied in the photon energy range of 1.3–3.0 eV by using a quartz-tungsten-halogen source. For PL studies, suspended samples were excited with a cw solid-state laser at a wavelength of 532 nm. A low laser power of $\sim$50 $\mu$W (on the sample) was used to avoid heating and PL saturation [26]. The PL QY of MoS$_2$ samples was calibrated by using rhodamine 6G molecules as the standard. For the photoconductivity studies, we prepared field-effect transistors of mono- and bilayer MoS$_2$ on silicon substrates covered by a 300-nm-thick oxide layer. Optical radiation of a well-defined frequency was selected from a supercontinuum laser source with a monochromator and was modulated with a mechanically thin layers of MoS$_2$ [Fig. 3(a)]. A PL QY on the order of $10^{-5}$–$10^{-6}$ was estimated for few-layer samples of $N = 2$–6; a value as high as $4 \times 10^{-3}$ was observed in the limit of monolayer thickness. In addition to the significant difference in the PL QY, the normalized PL spectra for mono- and few-layer samples are quite distinct from one another [Fig. 3(b)]. The PL spectrum of suspended monolayer samples consists of a single narrow feature of $\sim$50 meV width, centered at 1.90 eV. In contrast, few-layer samples display multiple emission peaks (labeled A, B, and I). Peak A coincides with the monolayer emission peak. It shifts to the red and broadens slightly with increasing $N$. Peak B lies about 150 meV above peak A. The broad feature I, which lies below peak A, systematically shifts to lower energies, approaching the indirect-gap energy of 1.29 eV, and becomes less prominent with increasing $N$ [Fig. 3(c)].

To understand the origin of these extraordinary PL properties, we compare the PL to the absorption spectrum of MoS$_2$. In the spectral range of 1.3–2.4 eV, absorption of bulk MoS$_2$ is dominated by two prominent transitions. They are known to arise from direct-gap transitions between the maxima of split valence bands ($\nu 1$ and $\nu 2$) and the minimum of the conduction band ($\nu 1$), all located at the $K$ point of the Brillouin zone (Fig. 1) [7,12–18]. The splitting arises from the combined effect of interlayer coupling and spin-orbit coupling. (The latter is not included in the band structure of Fig. 1 for simplicity.) The absorption spectrum of atomically thin layers of MoS$_2$ (normalized for thickness) was found to be largely unaltered, except for a slight blueshift of the resonances [Fig. 4(a)]. The monolayer PL peak A at 1.90 eV matches the lower absorption resonance in both its position and width. Therefore, we attribute the PL of monolayer MoS$_2$ to direct-gap luminescence. The result, combined with the high PL QY, also demonstrates the high quality of the suspended monolayer samples. For bilayer samples, the other hand, emission peaks A and B match the two absorption resonances and are assigned to direct-gap hot luminescence [27]. The weak feature between A and B could be impurity or defect luminescence and merits further investigation. The PL peak I at 1.59 eV ($\sim$ 300 meV below the direct-gap absorption peak), as we discuss below, is attributed to indirect-gap luminescence.

Probing of the indirect-gap optical transitions in atomically thin layers of MoS$_2$ is limited by the sensitivity of absorption measurements. In the case of weak exciton binding, as for MoS$_2$ at room temperature, the photoconductivity spectrum mimics the absorption spectrum. Therefore, we used photoconductivity spectroscopy to investigate the optical response below the direct gap of mono- and bilayer MoS$_2$ [12]. The photoconductivity spectra of these samples could be recorded with signals ranging over three decades in strength [Fig. 4(b)]. For bilayer MoS$_2$, the onset of photoconductivity occurs around 1.60 eV, coinciding with the PL peak I. The conductivity increases slowly with photon energy towards the direct band gap, as expected for an indirect-gap material. In contrast, for monolayer MoS$_2$, the photoconductivity near 1.8 eV increases.

![FIG. 2](image-url)

*(a) Representative optical image of mono- and few-layer MoS$_2$ crystals on a silicon substrate with etched holes of 1.0 and 1.5 $\mu$m in diameter. (b) PL image of the same samples. The PL QY is much enhanced for suspended regions of the monolayer samples, and the emission from the few-layer sample is too weak to be seen in this image.*
(from a negligible value) by about 3 orders of magnitude. This dramatic rise corresponds to the onset of optical absorption from the direct band edge [28].

In the following simplified treatment of the spectral dependence of the photoconductivity, we neglect both excitonic effects and the variation of matrix elements with energy, factors that should be included in a more comprehensive theory. The absorbance $A(\hbar \omega)$ at photon energy $\hbar \omega$ near a direct band edge of energy $E_g$ is then determined by the joint density of states. For a two-dimensional (2D) material like our atomically thin layers of MoS$_2$, this is described by a step function, $\Theta(\hbar \omega - E_g)$ [1,29]. After including a phenomenological broadening of 30 meV to account for finite temperature and scattering rates, we find that the photoconductivity spectrum of the monolayer samples can be fit well to this simple model [Fig. 4(b)].

This indicates that monolayer MoS$_2$ is indeed a direct-gap material [28]. On the other hand, the photoconductivity spectrum for bilayer MoS$_2$ cannot be described by the step-function response. We need to include the effect of an indirect transition. Near an indirect band edge, the corresponding absorbance can be represented by [1,29]

$$A(\hbar \omega) \propto \sum_k \left[ \frac{1+\exp(-\beta \hbar \omega/kT)}{1+\exp(-\beta \hbar \omega/kT)} \right] \hbar \omega - E'_g.$$ 

Here $E'_g$ and $\hbar \Omega_n$ denote, respectively, the indirect-gap energy and that of the $n$th phonon mode, and $kT$ is the thermal energy. By taking this term into account, the experimental bilayer MoS$_2$ spectrum can be fit well by an indirect transition at 1.6 eV, combined with a direct transition at 1.88 eV [Fig. 4(b)].

The indirect-direct-gap crossover as a function of MoS$_2$ thickness $N$ is the result of a significant upshift of $E'_g$ induced by perpendicular quantum confinement. To understand this, let us examine the electronic band structure of bulk MoS$_2$ (Fig. 1). The direct gap of $\sim 1.8$ eV occurs between $c_1$ and $v_1$ at the $K$ point of the Brillouin zone (transitions A) [3–5,7,12]. On the other hand, the maximum of $v_1$ and minimum of $c_1$ are located at the $\Gamma$ point and

![Figure 3](image-url) FIG. 3 (color online). (a) PL spectra for mono- and bilayer MoS$_2$ samples in the photon energy range from 1.3 to 2.2 eV. Inset: PL QY of thin layers for $N = 1–6$. (b) Normalized PL spectra by the intensity of peak A of thin layers of MoS$_2$ for $N = 1–6$. Feature I for $N = 4–6$ is magnified and the spectra are displaced for clarity. (c) Band-gap energy of thin layers of MoS$_2$, inferred from the energy of the PL feature I for $N = 2–6$ and from the energy of the PL peak A for $N = 1$. The dashed line represents the (indirect) band-gap energy of bulk MoS$_2$.

![Figure 4](image-url) FIG. 4 (color online). (a) Absorption spectra (left axis, normalized by $N$) and the corresponding PL spectra (right axis, normalized by the intensity of the peak A). The spectra are displaced along the vertical axis for clarity. (b) Photoconductivity spectra for mono- (red dots) and bilayer (green dots) samples [27]. The data are compared with the 2D model described in the text (blue lines). $E_g$ and $E'_g$ inferred from the PL measurements, indicated by arrows, are included for comparison.
along the Γ-K direction [3–6,8,11], respectively; they form an indirect gap of 1.29 eV (transition $I$) [20]. The out-of-plane mass $m_\perp$ for electrons and holes around the $K$ point far exceeds the free electron mass $m_0$; on the other hand, $m_\perp$ for holes around the $\Gamma$ point is estimated to be $\sim 0.4m_0$ and for electrons around the $c1$ minimum along the Γ-K direction to be $\sim 0.6m_0$. Thus, while decreasing $N$ leads to a large confinement-induced increase in the indirect gap $E_g^c$, the direct gap $E_g$ remains almost unchanged. The indirect-direct-gap crossover is reached in the limit of monolayer thickness.

To explain this result more precisely, we apply a zone-folding scheme to describe the electronic states of the ultrathin films [30,31]. For identical layers with only nearest-neighbor interlayer interactions, the 2D electronic structure of few-layer MoS$_2$ samples can be generated as a subset of the bulk electronic states with quantized out-of-plane momenta $k_\perp$, i.e., momenta lying in planes perpendicular to the Γ-A or $K$-$H$ directions. For monolayer MoS$_2$, the allowed plane passes directly through the $H$ and $A$ points. The $c1$ minimum and the $v1$ maximum of the resulting 2D bands both occur at the $H$ point (Fig. 1). Monolayer MoS$_2$ therefore becomes a direct-gap material. With increasing $N$, cuts with $k_\perp$ approaching the Γ-$K$ line are made. Then the $v1$ maximum and the $c1$ minimum occur at the $\Gamma$ point and along the Γ-K direction, respectively, as in the bulk material. Few-layer MoS$_2$ samples are, therefore, indirect-gap semiconductors. These findings agree with earlier *ab initio* calculations of few-layer MoS$_2$ [8,9]. Our result is distinct from zone-folding-induced pseudo-direct transitions in semiconductor superlattice structures [32], which have very low oscillator strength and PL QY.

The crossover from an indirect-gap material to a direct-gap material naturally accounts for the more than 10$^4$-fold enhancement of the luminescence QY observed in monolayer MoS$_2$. It is consistent with the observed absorption spectra that show little sensitivity to the layer number $N$ and a band gap that decreases with increasing $N$. The observed dependence of the band gap on $N$ is also in qualitative agreement with published density-functional theory calculations [8]. Monolayer MoS$_2$ constitutes the first atomically thin material that is an effective emitter of light. Its strong PL suggests its use for photostable markers and sensors that can be adapted to probe nanoscale dimensions. The controllability of the band gap may also be used to optimize the material’s use as a photocatalytic agent [19] and for photovoltaic applications. Our results highlight that the distinctive electronic properties of atomically thin layered materials are not restricted to graphene but extend to a broader group of van der Waals bonded solids.

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[22] Changes in the phonon spectra as a function of the layer thickness of MoS$_2$ have also been observed by using Raman spectroscopy: C. Lee et al., *ACS Nano* 4, 2695 (2010).

[23] Enhancement of PL QY in the MoS$_2$ monolayer samples has recently been reported in an independent study: A. Splendiani et al., *Nano Lett.* 10, 1271 (2010).


[28] The indirect-to-direct band-gap crossover is determined with a resolution on the order of the thermal energy $kT \sim 26$ meV.


