SPECIAL TOPIC

MoS₂ saturable absorber prepared by chemical vapor deposition method for nonlinear control in Q-switching fiber laser

To cite this article: Meng-Li Liu et al 2018 Chinese Phys. B 27 084211

View the article online for updates and enhancements.

Related content

al.

- <u>Large-area and highly crystalline MoSe2</u> for optical modulator Jinde Yin, Hao Chen, Wei Lu et al.
- CVD-grown MoSe2 with high modulation depth for ultrafast mode-locked erbiumdoped fiber laser
- Wenjun Liu, Mengli Liu, Yuyi OuYang et al.
- <u>Tungsten diselenide for mode-locked</u> erbium-doped fiber lasers with short pulse duration Wenjun Liu, Mengli Liu, Yuyi OuYang et

This content was downloaded from IP address 159.226.35.202 on 08/10/2018 at 09:25

SPECIAL TOPIC — Nanophotonics

MoS₂ saturable absorber prepared by chemical vapor deposition method for nonlinear control in Q-switching fiber laser *

Meng-Li Liu(刘孟丽)¹, Yu-Yi OuYang(欧阳毓一)¹, Huan-Ran Hou(侯焕然)¹, Ming Lei(雷鸣)¹, Wen-Jun Liu(刘文军)^{1,2,†}, and Zhi-Yi Wei(魏志义)^{2,‡}

¹State Key Laboratory of Information Photonics and Optical Communications, School of Science, Beijing University of Posts and Telecommunications, Beijing 100876, China ²Beijing National Laboratory for Condensed Matter Physics, Institute of Physics, Chinese Academy of Sciences, Beijing 100190, China

(Received 25 April 2018; revised manuscript received 20 May 2018; published online 10 July 2018)

Due to the remarkable carrier mobility and nonlinear characteristic, MoS₂ is considered to be a powerful competitor as an effective optical modulated material in fiber lasers. In this paper, the MoS₂ films are prepared by the chemical vapor deposition method to guarantee the high quality of the crystal lattice and uniform thickness. The transfer of the films to microfiber and the operation of gold plated films ensure there is no heat-resistant damage and anti-oxidation. The modulation depth of the prepared integrated microfiber- MoS_2 saturable absorber is 11.07%. When the microfiber- MoS_2 saturable absorber is used as a light modulator in the Q-switching fiber laser, the stable pulse train with a pulse duration of 888 ns at 1530.9 nm is obtained. The ultimate output power and pulse energy of output pulses are 18.8 mW and 88 nJ, respectively. The signal-to-noise ratio up to 60 dB indicates the good stability of the laser. This work demonstrates that the MoS₂ saturable absorber prepared by the chemical vapor deposition method can serve as an effective nonlinear control device for the Q-switching fiber laser.

Keywords: nonlinear optical materials, fiber laser, Q-switching

PACS: 42.70.Mp, 42.55.Wd, 42.60.Gd

DOI: 10.1088/1674-1056/27/8/084211

1. Introduction

The saturable absorber (SA), as a natural light modulation device, is the key part of the pulsed fiber laser in Q-switching and mode-locking. Semiconductor saturable absorber mirrors (SESAMs) are the most well-researched and mature SAs, and we attribute the success in application to the controllable absorption wavelength, saturation threshold, modulation depth, and relaxation time.^[1,2] However, the relatively high cost and narrow working bandwidth of SESAM have urged researchers to seek new alternatives for them. Then, the rise of graphene has meant two-dimensional (2D) materials are gradually being used more and more.^[3] Some of the most representative materials that can be used as SAs have been studied in depth.^[4,5] The unique zero band gap structure of graphene impels it to own the wider working wavelength than other 2D materials. The ultrafast recovery time and strong nonlinearity of graphene further prove that graphene can be a powerful candidate for SA.^[6-9] Topological insulators (TIs) are also interesting because of their large modulation depth and nonlinearity.^[10-13] The superiorities of TIs in light modulation contribute to their great potential in optoelectronics applications. Black phosphorus (BPs) maintain a direct band gap whether they are a bulk or layered structure. Their band gap is adjustable from 0.3 eV to 1.5 eV, which indicates that it is easy to make a breakthrough in the application of photon and photoelectron in the near infrared band.^[14-18] In addition to these classic materials, some novel 2D materials have also begun to arouse interest due to their unique electronic and optical characteristics. BP quantum dots (BPQDs), which are able to combine the quantum confinement and edge effects, exhibit excellent nonlinear optical response according to previous studies.^[19] Moreover, BPODs have been successfully used in mode-locked erbium-doped fiber laser (EDFL). MXene, as another kind of new member of the 2D material family, shows good conductivity, tunable bandgap, and ability to perform ion intercalation. It has reported that MXene performs optical manipulation in photonic applications, similar to or better than graphene.^[20]

In transition metal dichalcogenides (TMDs), a layer of M (Mo, W, etc.) atoms is sandwiched between two layers of X (S, Se, etc.) atoms, so the chemical formula of TMDs is generally written as MX_2 . The neighbor layers of TMDs are bonded with weak van der Waals force, which indicates layered nanosheets are easily separated from the sample.^[21-36] From previous work, the band gap of TMD varies with the number of lay-

[†]Corresponding author. E-mail: jungliu@bupt.edu.cn

[‡]Corresponding author. E-mail: zywei@iphy.ac.cn

^{*}Project supported by the National Natural Science Foundation of China (Grant No. 11674036), the Beijing Youth Top-notch Talent Support Program, China (Grant No. 2017000026833ZK08), the Fund of State Key Laboratory of Information Photonics and Optical Communications, Beijing University of Posts and Telecommunications, China (Grant Nos. IPOC2016ZT04 and IPOC2017ZZ05).

^{© 2018} Chinese Physical Society and IOP Publishing Ltd

ers, and so the consequent layer-dependent electronic and optical characteristics are attracting increasing attention.^[37–39] Among TMDs, MoS₂ is prominent considering its performance in photoelectrons. It is reported that the nonlinear optical response of MoS₂ is better than graphene.^[40,41] The few-layer MoS₂ unambiguously exhibits broadband absorption from previous work.^[42] Furthermore, it is found that the starting Q-switching operation threshold of MoS₂ is small under the same experimental conditions compared with those of the materials of the same kind.^[43] The relaxation time of MoS₂ is estimated at ~ 30 fs as mentioned in Ref. [44], which indicates that it has potential for fabricating the ultra-fast electronic devices.

There are various methods to be chosen in the fabrication of SA. The two most representative top-down methods are mechanical exfoliation (ME)^[45,46] with the aid of scotch tapes, and liquid-phase exfoliation (LPE)^[47,48] getting dispersant through ultrasonication and centrifugation. These two methods are convenient and low-cost, but the layer number of prepared nanosheets is random. By comparison, the three bottom-up methods, as the common methods, have better performance in the control of the layer number and the quality of the sample, they are the pulsed laser deposition (PLD) method,^[49,50] chemical vapor deposition (CVD) method,^[51,52] and magnetron sputtering deposition (MSD) method,^[53,54] respectively. From previous reports, the layer number of nanosheets prepared by the CVD method tends to be controlled more accurately through adjusting the reaction parameters.

Here in this paper, the MoS₂ produced through the CVD method is moved to the waist of the microfiber to form MoS₂ SA. Finally, the effective area of the taper fiber is fully covered with gold to prevent it from being oxidized. By means of the interaction between the evanescent field of light and the material, the SA is able to avoid some thermal damage.^[55] Furthermore, the long reaction length promotes the MoS₂ SA to fully exhibit the optical nonlinearity. The modulation depth (MD) and saturable intensity of MoS2 SA are measured to be 11.07% and 1.871 MW/cm², respectively. The proposed MoS₂ based Q-switching EDFL operates at 1530.9 nm. The corresponding pulse duration and maximum output power are 888 ns and 18.8 mW, respectively. The signal-to-noise ratio (SNR) (up to 60 dB) indicates the stability of the laser. The experimental results further prove the favorable optical modulation ability of MoS₂ SA.

2. Preparation and characterization of MoS₂ SA

The MoS_2 nanosheets were grown on the ultrathin quartz plate by the CVD method. Before the synthesis, the sulfur powder of 0.5 g was placed in the reaction chamber, and the MoO_3 powder of 0.5 g was placed in the heating center, which was at the downstream of sulfur powder. The ultrathin quartz plate was placed at the downstream of the heating center, which was 20 cm away from MoO₃ powder. During the synthesis, the pressure of the chamber was set to be 0.1 Torr (1 Torr = 1.33322×10^2 Pa). Under the drive of Ar flow gas, MoO₃ and sulfur vapors were capable of mixing and reacting fully, as the product was brought to the substrate. Because temperature is a key factor for determining the quality of the finished product, the temperature of the chamber was rising at a constant speed of 25 °C/min. The reaction chamber was heated up to 550 °C and maintained this temperature for 0.5 h, then the reaction was finished.

When the chamber cooled down to room temperature, ultrathin quartz plate attached with MoS₂ film was achieved. To prevent the lattice structure of MoS₂ film from destructing during transfer, the polymethyl methacrylate (PMMA) assisted method was adopted. Before the transfer, the PMMA was spin coated on a large scale over the MoS₂ film. The MoS₂ PMMA was able to be easily separated from ultrathin quartz plate after the open-air drying of the sample. To obtain the pure MoS₂ film, the MoS₂PMMA was etched in acetone. Then the pure MoS₂ film was soaked in deionized water, which could not only remove the remnants, but also make the film float smoothly with the help of water tension. Finally, the MoS₂ film was moved to the conical area of microfiber (SMF-28e), which was observed by using a microscope. The measured diameter of the tapered waist was 12 µm, and the effective contact length between the MoS₂ film and fiber was about 2 mm. After the volatilization of superfluous deionized water, the MoS₂ film closely adhered to the lateral wall of tapered fiber due to van der Waals interaction. Finally, the effective area of the microfiber was fully covered with gold to avoid rapidly oxidizing the material.



Fig. 1. (color online) (a) Surface morphology image of MoS_2 film; (b) altitude difference of the surface.

With the aid of the atomic force microscope (AFM), an important device for the nano-scale measurement, we obtained the morphology of the MoS_2 surface without damaging the material. Figure 1(a) shows the particles of the material are arranged closely and orderly, which indicates the MoS_2 films prepared by the CVD method are of great uniformity and crystallinity. The comparison of surface height among different regions on the MoS_2 films shows the thickness of MoS_2 film was 30 nm as indicated in Fig. 1(b), which corresponds to 41–42 layers in structure.



Fig. 2. (color online) Raman spectra of prepared MoS₂ films.

Raman spectroscopy is considered to be an effective method of determining the specific vibration modes of the material. In Fig. 2, two distinct characteristic peaks are located at 378.9 cm⁻¹ and 403.6 cm⁻¹. According to previous research, the mode at 378.9 cm⁻¹ comes from the inplane vibration which belongs to Mo and S atoms, while the mode at 403.6 cm⁻¹ comes from the out-of-plane vibration which belongs to S atoms.^[56–58] The frequency difference between two peaks illustrates the multilayer structure of prepared MoS₂ film.



Fig. 3. (color online) Schematic diagram of power-dependent feature measurement.

The schematic diagram of power-dependent feature measurement is shown in Fig. 3. The light source was a modelocked fiber laser operating at 1561 nm, of which the pulse duration and the repetition rate were 200 fs and 67 MHz. A variable fiber optic attenuator (VOA) was used to manually adjust the signal attenuation when the optical signal passes through the device. By employing OC, the optical signal could be divided into two parts: one passed through MoS_2 SA to measure its saturated absorptive properties, and the other passed through SMF 28e for comparison. The optical signals at both ends were captured and recorded simultaneously by a power meter. Experimental data could be fitted by

$$\alpha(I) = \frac{\alpha_{\rm s}}{1 + I/I_{\rm sat}} + \alpha_{\rm ns}$$

where α_s , α_{ns} , and I_{sat} are the modulation depth, nonsaturable absorption, and saturation intensity. After the fitting of experimental data in Fig. 4, the saturable intensity and MD of MoS₂ SA are 1.871 MW/cm² and 11.07%, respectively.



Fig. 4. (color online) Intensity-dependent transmission of MoS_2 SA, showing its nonlinear absorber characteristics.

In Table 1, the nonlinear parameters of MoS₂ SAs obtained with different fabrication methods are listed to compare the advantages of CVD-MoS₂. By comparison, it can be found that CVD-MoS₂ generally has a large modulation depth, because the materials prepared by the CVD method have better uniformity, which is beneficial to the large modulation depth of SA. In addition, we find that the saturation intensity of CVD-MoS₂ is generally small, which indicates that the laser based on CVD-MoS₂ SA has a lower starting Q-switching operation threshold.

Material	Fabrication	MD/%	Saturation intensity	Non-saturable loss/%	Ref.
MoS ₂	Scotch-tape micromechanical cleavage technique method	9.70	_	28.80	[59]
MoS_2	Hydrothermal intercalation/exfoliation approach	10.47	-	-	[40]
MoS_2	LPE	4	_	55.90	[<mark>60</mark>]
MoS_2	LPE	2	10 MW/cm ²	48.56	[61]
MoS_2	LPE	2.15	129.4 MW/cm ²	63.10	[43]
MoS_2	LPE	6.30	1.6 MW/cm ²	18	[62]
MoS_2	LPE	1.60	13 MW/cm ²	54.80	[<mark>63</mark>]
MoS ₂	PLD	9	27 MW/cm ²	40.50	[<mark>64</mark>]
MoS ₂	CVD	35.40	0.34 MW/cm ²	34.10	[65]
MoS ₂	CVD	28.50	0.55 MW/cm ²	35	[66]
MoS ₂	CVD	11.07	1.871 MW/cm ²	77.61	this work
		094211 2			

 Table 1. Nonlinear parameters of MoS₂ SAs obtained with different fabrication methods.

3. Q-switching EDFL employed with MoS₂ SA

An all-fiber ring cavity in Fig. 5 was used in the experiment. Optical devices were coupled into the laser cavity in the order corresponding to the direction of propagation of light. The pump which operated at 976 nm had the maximum allowed output power of 630 mW. The 980/1550 nm WDM successfully coupled the pump light into the ring cavity for exciting the erbium-doped fiber (EDF). The group velocity dispersion (GVD) of EDF (Liekki 110-4/125) was 12 ps²/km at 1550 nm. The intra-cavity polarization and birefringence were mainly optimized by polarization controller (PC). The primary role of an isolator (ISO) is to guarantee unidirectional optical transmission in the cavity to avoid damaging the device, caused by redundant reverse light. The MoS₂ SA was placed between the ISO and WDM. The 20:80 optical coupler (OC) was adopted in the experiment, 20% of which was exported to monitor the real-time experimental data. The monitoring instruments used in the experiment mainly included the RF spectrum analyzer (Agilent E4402B), oscilloscope (Tektronix DPO3054), and optical spectrum analyzer (Yokogawa AQ6370C).

The stable Q-switching pulses occur at 147 mW with the increase of pump power. Meanwhile, the repetition rates of

the output pulses with different powers are recorded and exhibited in Fig. 6(a). The tunable range of frequency is 92 kHz– 212 kHz. When the pump power is increased to 630 mW, the shortest pulse duration of 888 ns is obtained as shown in Fig. 6(b). The accomplished Q-switching EDFL operates at 1530.9 nm, the corresponding spectral width is 3.14 nm. For a few-hours laser operation, the time-section data of the spectrum is shown in Fig. 6(c). The almost invariable spectral shape indicates the stability of the laser. The RF spectrum of Q-switching EDFL is provided in Fig. 6(d). The SNR of the fundamental frequency increases up to 60 dB, which indicates that the operative condition of the laser is relatively stable. Figure 6(d) shows that the downward trend of frequency multiplication is uniform, which further illustrates the stability of the laser.



Fig. 5. (color online) Experimental illustrative diagram of Q-switching EDFL employed with MoS₂ SA.



Fig. 6. (color online) Experimental date of Q-switching EDFL employed with MoS_2 SA, showing (a) Q-switching pulse trains at different powers, (b) single pulse waveform of Q-switching EDFL, (c) spectra at different times, and (d) RF spectrum of Q-switching EDFL.

According to the experimental monitoring data at different powers, the variation tendency of pulse duration and frequency rate with input power are shown in Fig. 7(a). In the prophase of the power growth, the pulse duration changes more sharply. However, in the later stage of power growth, the pulse duration tends to be stable. This illustrates that MoS₂ SA is close to saturation at high power. With the increase of power, the repetition rate is almost increased uniformly. Similarly, we record and calculate the corresponding output power and pulse energy of the Q-switched pulse train varying with pump power in Fig. 7(b). On the whole, not only output power but also pulse energy increase evenly with input power. The ultimate output power and pulse energy are obtained to be 18.8 mW and 88 nJ, respectively. The maximum damage threshold is about 19.5 mJ/cm².

Performance comparisons of Q-switched EDFL based on different SAs are shown in Table 2. By comparison, the CVD-MoS₂ has certain potentials and advantages in realizing short pulse duration. Moreover, the SNR of most Q-switched EDFL is below 50 dB. The SNR of Q-switched EDFL based on CVD-MoS₂ increases up to 60 dB, which indicates the stability of operation. These results show that the microfiber-MoS₂ SA prepared by the CVD method is able to support stable Qswitching operation, which means it can serve as an impactful nonlinear control device for Q-switching EDFL.



Fig. 7. (color online) (a) Variation of pulse duration and frequency rate with pump power, and (b) variation of output power and pulse energy with pump power.

Meterials	$\alpha_{\rm s}/\%$	$ au/\mu s$	Frequency/kHz	Energy/nJ	SNR/dB	Output power/mW	Ref.
MoS ₂	4	3.53	72.74-86.39	74.93	51.6	6.47	[60]
WS_2	4.48	0.16	91-318	54.4	40	17.3	[67]
WSe ₂	3.02	4.1	46.3-85.4	484.8	41.9	3.16	[43]
MoSe ₂	-	30.4	16.9-32.8	57.9	-	1.9	[68]
Bi ₂ Se ₃	4.3	1.9	495-940	23.8	50	22.35	[69]
BP	18.55	10.32	6.983-15.78	94.3	45	1.5	[45]
MoS_2	11.07	0.888	92–212	88	60	18.8	this work

Table 2. Performance comparisons of Q-switched EDFL based on different SAs.

4. Conclusions

By combining the CVD method with the PMMA assisted method, the MoS₂ SA based on microfiber is successfully produced. On the one hand, the films prepared by the CVD method are of good uniformity. On the other hand, the gold-plated SA based on microfiber has some advantages in preventing heat-resistant damage, and anti-oxidation. The saturate intensity and MD of MoS₂ SA are 1.871 MW/cm² and 11.07%, respectively. Moreover, the obtained MoS₂ SA is successfully applied to the Q-switching EDFL. The shortest pulse duration of laser is 888 ns, and the tunable range of frequency is 92 kHz–212 kHz. The ultimate output power and pulse energy are 18.8 mW and 88 nJ, respectively. The SNR of laser increases up to 60 dB, which illustrates the stability of the operation. These results show that the microfiber-MoS₂ SA prepared by the CVD method is able to support stable Q-

switching pulse output, and it can serve as an impactful nonlinear control device for Q-switching EDFL.

References

- [1] Keller U 2003 Nature 424 831
- [2] Okhotnikov O, Grudinin A and Pessa M 2004 New J. Phys. 6 177
- [3] Bao Q L, Zhang H, Wang Y, Ni Z H, Yan Y L, Shen Z X, Loh K P and Tang D Y 2009 Adv. Funct. Mater. 19 3077
- [4] Liu W J, Liu M L, Yin J D, Chen H, Lu W, Fang S B, Teng H, Lei M, Yan P G and Wei Z Y 2018 *Nanoscale* 10 7971
- [5] Liu W J, Liu M L, OuYang Y Y, Hou H R, Ma G L, Lei M and Wei Z Y 2018 Nanotechnology 29 174002
- [6] Zhang H, Bao Q L, Tang D Y, Zhao L M and Loh K P 2009 Appl. Phys. Lett. 95 141103
- [7] Sun Z P, Hasan T, Torrisi F, Popa D, Privitera G, Wang F Q, Bonaccorso F, Basko D M and Ferrari A C 2010 ACS Nano 4 803
- [8] Luo A P, Zhu P F, Liu H, Zheng X W, Zhao N, Liu M, Cui H, Luo Z C and Xu W C 2014 Opt. Express 22 27019
- [9] Sotor J, Bogusławski J, Martynkien T, Mergo P, Krajewska A, Przewłoka A, StrupiŃski W and SoboŃ G 2017 Opt. Lett. 42 1592

- [10] Luo Z Q, Liu C, Huang Y Z, Wu D D, Wu J Y, Xu H Y, Cai Z P, Lin Z Q, Sun L P and Weng J 2014 IEEE J. Sel. Top. Quantum Electron. 20 0902708
- [11] Sotor J, Sobon G, Grodecki K and Abramski K M 2014 Appl. Phys. Lett. 104 251112
- [12] Liu H, Zheng X W, Liu M, Zhao N, Luo A P, Luo Z C, Xu W C, Zhang H, Zhao C J and Wen S C 2014 Opt. Express 22 6868
- [13] Liu W J, Pang L H, Han H N, Tian W L, Chen H, Lei M, Yan P G and Wei Z Y 2016 Sci. Rep. 6 19997
- [14] Sotor J, Sobon G, Macherzynski W, Paletko P and Abramski K M 2015 Appl. Phys. Lett. 107 051108
- [15] Hu G H, Albrow-Owen T, Jin X X, Ali A, Hu Y W, Howe R C T, Shehzad K, Yang Z Y, Zhu X K, Woodward R I, Wu T C, Jussila H, Wu J B, Peng P, Tan P H, Sun Z P, Kelleher E J R, Zhang M, Xu Y and Hasan T 2017 *Nat. Commun.* 8 278
- [16] Li J F, Luo H Y, Zhai B, Lu R G, Guo Z N, Zhang H and Liu Y 2016 Sci. Rep. 6 30361
- [17] Liu S C, Zhang Y N, Li L, Wang Y G, Lv R D, Wang X, Chen Z D and Wei L L 2018 Appl. Opt. 57 1292
- [18] Yun L 2017 Opt. Express 25 32380
- [19] Xu Y H, Wang Z T, Guo Z N, Huang H, Xiao Q L and Zhang H 2016 Adv. Opt. Mater. 4 1223
- [20] Jiang X T, Liu S X, Liang W Y, Luo S J, He Z L, Ge Y Q, Wang H D, Cao R, Zhang F, Wen Q, Li J Q, Bao Q L, Fan D Y and Zhang H 2018 *Laser Photon. Rev.* 12 1700229
- [21] Li W Y, OuYang Y Y, Ma G L, Liu M L and Liu W J 2018 Laser Phys. 28 055104
- [22] Liu W J, Liu M L, Lei M, Fang S B and Wei Z Y 2018 IEEE J. Sel. Top. Quantum Electron. 24 0901005
- [23] Yang C Y, Li W Y, Yu W T, Liu M L, Zhang Y J, Ma G L, Lei M and Liu W J 2018 Nonlinear Dyn. 92 203
- [24] Zhang M, Howe R C T, Woodward R I, Kelleher E J R, Torrisi F, Hu G H, Popov S V, Taylor J R and Hasan T 2015 Nano Res. 8 1522
- [25] Pumera M, Sofer Z and Ambrosi A 2014 J. Mater. Chem. A 2 8981
- [26] Yan P G, Liu A J, Chen Y S, Chen H, Ruan S C, Guo C Y, Chen S F, Li I L, Yang H P, Hu J G and Cao G Z 2015 Opt. Mater. Express 5 479
- [27] Li W Y, Ma G L, Yu W T, Zhang Y J, Liu M L, Yang C Y and Liu W J 2018 Chin. Phys. B 27 030504
- [28] Liu M L, Liu W J, Yan P G, Fang S B, Teng H and Wei Z Y 2018 Chin. Opt. Lett. 16 020007
- [29] Yu W T, Yang C Y, Liu M L, Zhang Y J and Liu W J 2018 Optik 159 21
- [30] Huang X, Zeng Z Y and Zhang H 2013 Chem. Soc. Rev. 42 1934
- [31] Wang J T, Jiang Z K, Chen H, Li J R, Yin J D, Wang J Z, He T C, Yan P G and Ruan S C 2017 *Opt. Lett.* 42 5010
- [32] Liu W J, Pang L H, Han H N, Shen Z W, Lei M, Teng H and Wei Z Y 2016 Photon. Res. 4 111
- [33] Zhang M, Hu G H, Hu G Q, Howe R C T, Chen L, Zheng Z and Hasan T 2015 Sci. Rep. 5 17482
- [34] Liu W J, Zhu Y N, Liu M L, Wen B, Fang S B, Teng H, Lei M, Liu L M and Wei Z Y 2018 Photon. Res. 6 220
- [35] Liu M L, Liu W J, Pang L H, Teng H, Fang S B and Wei Z Y 2018 Opt. Commun. 406 72
- [36] Liu W J, Yang C Y, Liu M L, Yu W T, Zhang Y J and Lei M 2017 *Phys. Rev. E* 96 042201
- [37] Mao D, She X Y, Du B B, Yang D X, Zhang W D, Song K, Cui X Q, Jiang B Q, Peng T and Zhao J L 2016 Sci. Rep. 6 23583
- [38] Wang Q H, Kalantar-Zadeh K, Kis A, Coleman J N and Strano M S 2012 Nat. Nanotechnol. 7 699
- [39] Britnell L, Ribeiro R M, Eckmann A, Jalil R, Belle B D, Mishchenko A, Kim Y J, Gorbachev R V, Georgiou T, Morozov S V, Grigorenko A N, Geim A K, Casiraghi C, Castro Neto A H and Novoselov K S 2013 *Science* 340 1311

- [40] Du J, Wang Q K, Jiang G B, Xu C W, Zhao C J, Xiang Y J, Chen Y, Wen S C and Zhang H 2015 Sci. Rep. 4 6346
- [41] Zhang H, Lu S B, Zheng J, Du J, Wen S C, Tang D Y and Loh K P 2014 Opt. Express 22 7249
- [42] Wang S X, Yu H H, Zhang H J, Wang A Z, Zhao M W, Chen Y X, Mei L M and Wang J Y 2014 Adv. Mater. 26 3538
- [43] Chen B H, Zhang X Y, Wu K, Wang H, Wang J and Chen J P 2015 Opt. Express 23 26723
- [44] Wang K P, Wang Jun, Fan J T, Lotya M, O'Neill A, Fox D, Feng Y Y, Zhang X Y, Jiang B X, Zhao Q Z, Zhang H Z, Coleman J N, Zhang L and Blau W J 2013 ACS Nano 7 9260
- [45] Chen Y, Jiang G B, Chen S Q, Guo Z N, Yu X F, Zhao C J, Zhang H, Bao Q L, Wen S C, Tang D Y and Fan D Y 2015 Opt. Express 23 12823
- [46] Aiub E J, Steinberg D, de Souza E A T and Saito L A M 2017 Opt. Express 25 10546
- [47] Wu K, Zhang X Y, Wang J, Li X and Chen J P 2015 Opt. Express 23 11453
- [48] Koo J, Park J, Lee J, Jhon Y M and Lee J H 2016 Opt. Express 24 10575
- [49] Guo B, Yao Y, Yan P G, Xu K, Liu J J, Wang S G and Li Yuan 2016 IEEE Photon. Technol. Lett. 28 323
- [50] Liu W J, Pang L H, Han H N, Liu M L, Lei M, Fang S B, Teng H and Wei Z Y 2017 Opt. Express 25 2950
- [51] Yan P G, Chen H, Yin J D, Xu Z H, Li J R, Jiang Z K, Zhang W F, Wang J Z, Li I L, Sun Z and Ruan S C 2017 Nanoscale 9 1871
- [52] Zhao L M, Tang D Y, Zhang H, Wu X, Bao Q L and Loh K P 2010 Opt. Lett. 35 3622
- [53] Chen H, Chen Y S, Yin J D, Zhang X J, Guo T and Yan P G 2016 Opt. Express 24 16287
- [54] Yan P G, Lin R Y, Ruan S C, Liu A J and Chen H 2015 Opt. Express 23 154
- [55] Wu K, Chen B H, Zhang X Y, Zhang S F, Guo C S, Li C, Xiao P S, Wang J, Zhou L J, Zou W W and Chen J P 2018 Opt. Commun. 406 214
- [56] Li H, Zhang Q, Yap C C R, Tay B K, Edwin T H T, Olivier A and Baillargeat D 2012 Adv. Funct. Mater. 22 1385
- [57] Zhao Y Y, Luo X, Li H, Zhang J, Araujo P T, Gan C K, Wu J, Zhang H, Quek S Y and Dresselhaus M S 2013 Nano Lett. 13 1007
- [58] Lee C, Yan H, Brus L E, Heinz T F, Hone J and Ryu S 2010 ACS Nano 4 2695
- [59] Lou F, Zhao R W, He J L, Jia Z T, Su Xi C, Wang Z W, Hou J and Zhang B T 2015 Photon. Res. 3 A25
- [60] Luo Z C, Wang F Z, Liu H, Liu M, Tang R, Luo A P and Xu W C 2016 Opt. Eng. 55 081308
- [61] Huang Y Z, Luo Z Q, Li Y Y, Zhong M, Xu B, Che K J, Xu H Y, Cai Z P, Peng J and Weng J 2014 Opt. Express 22 25258
- [62] Woodward R I, Kelleher E J R, Howe R C T, Hu G, Torrisi F, Hasan T, Popov S V and Taylor J R 2014 Opt. Express 22 31113
- [63] Luo Z Q, Huang Y Z, Zhong M, Li Y Y, Wu J Y, Xu B, Xu H Y, Cai Z P, Peng J and Weng 2014 J. Lightwave Technol. 32 4077
- [64] Ren J, Wang S X, Cheng Z C, Yu H H, Zhang H J, Chen Y X, Mei L M and Wang P 2015 Opt. Express 23 5607
- [65] Xia H D, Li H P, Lan C Y, Li C, Zhang X X, Zhang S J and Liu Y 2014 Opt. Express 22 17341
- [66] Xia H D, Li H P, Lan C Y, Li C, Du J B, Zhang S J and LiuY 2015 Photon. Res. 3 A92
- [67] Chen H, Li L, Ruan S C, Guo T and Yan P G 2016 Opt. Eng. 55 081318
- [68] Ahmad H, Suthaskumar M, Tiu Z C, Zarei A and Harun S W 2016 Opt. Laser Technol. 79 20
- [69] Yu Z H, Song Y R, Tian J R, Dou Z Y, Guoyu H Y, Li K X, Li H W and Zhang X P 2014 Opt. Express 22 11508