WS$_2$ mode-locked ultrafast fiber laser

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Graphene-like two dimensional materials, such as WS$_2$ and MoS$_2$, are highly anisotropic layered compounds that have attracted growing interest from basic research to practical applications. Similar with MoS$_2$, few-layer WS$_2$ has remarkable physical properties. Here, we demonstrate for the first time that WS$_2$ nanosheets exhibit ultrafast nonlinear saturable absorption property and high optical damage threshold. Soliton mode-locking operations are achieved separately in an erbium-doped fiber laser using two types of WS$_2$-based saturable absorbers, one of which is fabricated by depositing WS$_2$ nanosheets on a D-shaped fiber, while the other is synthesized by mixing WS$_2$ solution with polyvinyl alcohol, and then evaporating them on a substrate. At the maximum pump power of 600 mW, two saturable absorbers can work stably at mode-locking state without damage, indicating that few-layer WS$_2$ is a promising high-power flexible saturable absorber for ultrafast optics. Numerous applications may benefit from the ultrafast nonlinear features of WS$_2$ nanosheets, such as high-power pulsed laser, materials processing, and frequency comb spectroscopy.

Since the single- or few-layer graphene was fabricated by using mechanical exfoliation method, it has been in-depth investigated and widely used in various fields, including photon detection, graphene plasmonics, ultrafast optics, and quantum electrodynamics$^{1-7}$. Research on graphene also opened up a door to new two-dimensional (2D) nanomaterials in which strong covalent bonds in layers and weak van der Waals interaction between layers exist. Following the similar approach on graphene study, researchers have developed various graphene analogues of layered inorganic materials comprising stacked atomic or molecular layers, such as topological insulator$^{6-8}$, molybdenum disulfide (MoS$_2$)$^9$, tungsten disulphide (WS$_2$)$^{10}$, etc. Among them, topological insulator nanosheets are proven to exhibit saturable absorption property$^{7,8}$, in which the optical absorbance decreases with the enhancement of the incident laser intensity and becomes saturated above a certain threshold. Graphene$^{11,12}$ and topological insulator$^{7,8,13}$ based saturable absorbers (SAs) possess inherent features of broad response as well as high flexibility, and have been broadly used in Q-switching and mode-locking lasers. Bonaccorso et al. summarized graphene and other 2D crystals based ultrafast photonics, from solution processing, fabrication of SAs, to their applications in ultrafast lasers$^{14}$.

The bulk MoS$_2$ and WS$_2$ are indirect semiconductors, which consist of hexagonal layers of metal atoms sandwiched between two layers of chalcogen atoms$^{15}$. Like graphene, these layered materials can be exfoliated into thin film to explore their exotic features for practical applications. Various methods, such as mechanical exfoliation$^{16}$, liquid exfoliation$^{17}$, hydrothermal intercalation/exfoliation$^{18}$, chemical vapor deposition$^{19}$, as well as two-step expansion and intercalation$^{20}$, have been developed to synthesize few-layer MoS$_2$ and WS$_2$ nanosheets. Compared with their bulk counterparts, the monolayer MoS$_2$ and WS$_2$ possess a direct bandgap and show significant improvement in photoluminescence quantum efficiency attributing to the 2D confinement of electron motion and the absence of interlayer perturbation$^{21,22}$. Recently, a great deal of research interest is focused on the nonlinear absorption property of MoS$_2$ nanosheets$^{23-25}$. For example, K. Wang et al. demonstrated that MoS$_2$ nanosheets exhibited significant saturable absorption with a relaxation of 30 fs and had better saturable absorption than that of graphene for femtosecond pulse at 800 nm$^{25}$. They concluded that the ultrafast saturable absorption of the MoS$_2$ nanosheets originated from one-photon induced free-carrier absorption. After that, S. Wang et al. showed that when the ratio between Mo and S ions was increased from 2 to 2.09, the MoS$_2$ bandgap can be reduced from 1.08 to 0.08 eV (corresponding to an absorption wavelength of 15.4 μm). By introducing suitable defects, they have realized a broadband MoS$_2$ SA that has been used in Q-switched lasers operating at 1.06, 1.42, and 2.1 μm$^{25}$. H. Zhang et al. systematically investigated the nonlinearity of MoS$_2$ nanosheets using open-aperture Z-scan and balanced-detector measurement techniques, and demonstrated a yttrium-doped fiber laser passively mode locked with a MoS$_2$ SA$^{26}$. They infer that the broadband absorption property may arise from the coexistence of semiconducting and metallic states in the layered 2D materials. Currently, H. Liu et al. and H.
Xia et al. have realized femtosecond and picosecond pulses in erbium-doped fiber (EDF) lasers mode locked by MoS$_2$ SAs, respectively$^{25,26}$. R. Khazaeizhad et al. have realized passive mode locking in anomalous- and normal-dispersion regimes using a multilayer MoS$_2$ thin film coated on D-shaped fiber$^{27}$. The aforementioned investigations are mainly focused on MoS$_2$, while that of WS$_2$ are not stressed and its applications remain untapped. Whether WS$_2$ exhibits saturable absorption property as that of graphene or MoS$_2$? How to fabricate WS$_2$-based mode locker to realize ultrafast pulse emission? In this paper, we focus on these important issues.

Here, we demonstrate that WS$_2$ nanosheets exhibit nonlinear saturable absorption property and ultra-high optical damage threshold. Soliton mode locking is achieved separately in a fiber laser by using two different types of WS$_2$ SAs, one of which is fabricated by depositing WS$_2$ on D-shaped fiber, while the other is synthesized by mixing WS$_2$ solution with polyvinyl alcohol (PVA), and then evaporating them on a substrate. At the maximum pump power of 600 mW, two SAs can work stably without damage in the mode-locked fiber laser, indicating that WS$_2$-based SAs possess ultra-high damage threshold. Our results demonstrate that WS$_2$ nanosheets exhibit saturable absorption property at 1.55 μm and can work as a high-power mode locker for ultrafast fiber lasers.

**Results**

**Preparation and characterization of WS$_2$-based SAs.** The liquid exfoliation is a simple and effective method for preparing high-quality 2D nanostructures$^{17,22}$. In this experiment, the solvent for dispersing WS$_2$ flakes is fabricated by mixing ethanol and water at the volume ratio of 35:75. The initial dispersions are treated for 120 min by a high-power ultrasonic cleaner. After sonication, the

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**Figure 1** | (a) Image of WS$_2$ dispersion; (b) SEM image, and (c) AFM image, (d) Raman spectrum of WS$_2$ nanosheets; (e) linear transmission spectrum of the WS$_2$-PVA film in comparison with pure PVA film; (f) side profile of WS$_2$-PVA film. The thickness of the film is measured as 36.31 μm.
dispersions are allowed to settle for several hours. To remove large agglomeration, the WS₂ dispersions are centrifuged at 3000 rpm for 30 min, and the upper supernatant is collected. The concentration of WS₂ nanosheets in the solvent is 0.1 mg/ml. As shown in Fig. 1(a), the typical dispersions show a faint yellow-green color, and are stable over tens of days. Figure 1(b) shows the scanning electron microscopy (SEM) image of WS₂ nanosheets on silicon slice. The width and length of WS₂ nanosheets are in the range of 50 to 500 nm. By using an atomic force microscope (AFM), we find that the thickness of nanosheets ranges from 2 to 20 nm. Figure 1(c) shows three typical AFM images of WS₂ nanosheets. The deposited WS₂ nanosheets are further characterized by Raman spectroscopy using Ar laser at 514 nm, as depicted in Fig. 1(d). The characteristic bands at 350.8 and 420.7 cm⁻¹ on the Raman spectrum are assigned to the in-plane (E₂g) and out-of-plane (A₁g) vibrational modes of WS₂, in agreement with the earlier findings. Figure 1(e) plots the linear transmission spectrum of the WS₂-PVA film in comparison with pure PVA film, which is measured by a spectrometer (Hitachi UV4100). The dip at 630 nm on the transmission spectrum could be attributed to the direct gap transition. The side profile of WS₂-PVA film is measured with a SEM, and the thickness of the film is given as 36.31 μm, as shown in Fig. 1(f). Analysis of SEM profile, AFM image, Raman spectrum, and linear transmission suggests the presence of WS₂ nanosheets in the sample.

In our experiment, two different WS₂ SAs are used to achieve passive mode locking. The first SA is realized by nonlinear interaction of WS₂ nanosheets with the evanescent field of light in a D-shaped fiber. Here, the D-shaped fiber is prepared by polishing a single-mode fiber (SMF) after holding the fiber with an arcuate block. During polishing, the insertion loss that indicates the space between the fiber core and the polished surface is monitored by an optical power meter. The WS₂ nanosheets are adhered on the D-shaped fiber by using optical deposition method. First, the WS₂ solution is dropped on a D-shaped fiber that is fixed at a quartz plate. Then, a 10 mW continuous wave centered at 1.55 μm is coupled into the D-shaped fiber and the optical deposition process starts. An optical power meter is used to evaluate the deposition depth real-timely. This approach is similar to the carbon nanotube mode locker via evanescent field interaction in D-shaped fibers or topological insulator mode locker based on tapered fibers.

The microscope images of the D-shaped fiber coated with WS₂ are shown in Fig. 2(a). The upward inset and middle images show the D-shaped fiber before and after injecting 632.8 nm laser, and the downward image shows the zoom-in image of D-shaped fiber. The insertion loss of the integrated D-shaped fiber is measured to be 0.8 dB with a continuous laser. The nonlinear saturable absorption of D-shaped fiber SA is measured by a power-dependent transmission technique based on a balanced twin-detector measurement system, which has been elaborated in Ref. 24. The illumination pulse is delivered by a homemade passively mode-locked fiber laser (repetition rate: 25 MHz, pulse duration: 400 fs, central wavelength: 1.55 μm). The power-dependent transmittance T is fitted by \( T = A \exp\left(-\Delta T/(1 + I/I_{sat})\right) \), where A is the normalization constant, \( \Delta T \) is the absolute modulation depth, I is the incident intensity, and \( I_{sat} \) is the saturation intensity. As shown in Fig. 2(b), the D-shaped fiber SA exhibits saturable absorption property that the transmittance increases with light intensity, and the modulation depth is measured as 0.95%. The saturable intensity is about 0.6 GW/cm², which is much larger than that of the graphene or carbon nanotube.

The second SA is based on WS₂-PVA film, and the fabrication process can be described as follows. The as-prepared WS₂ solution is ultra-sonicated for 1-hour using a 300 W ultrasonic cleaner. Then, 5 wt% aqueous PVA solution and the WS₂ solution are mixed at the
volume ratio of 1:2 by a magnetic stirrer for 5 hours. The final mixture is dropped on a substrate. Slow evaporation under ambient temperature and pressure results in a PVA-composite film. The side profile and thickness of the film are shown in Fig. 1(f). Figure 2(c) shows the photograph of WS2-PVA film that is coated on the facet of the fiber connector. In this case, the modulation depth and saturable power are given as about 1.8% and 0.75 GW/cm² by using the same measuring technique. Compared with dropping dispersions on a quartz plate and optical deposition, the filmy polymer method would be more favorable to fabricate SA due to its superiority of low cost, flexibility, and controllability. During the experiment, we do not observe any other nonlinear response from fiber device or pure PVA film, indicating that the saturable absorption originates from WS2 nanosheets alone.

**Setup of WS2 mode locked fiber laser.** Fiber lasers exhibit inherent advantages of high gain coefficient, wide operating wavelength range, excellent heat dissipation, and robust mode confinement and thus offer an attractive platform for testing the nonlinear absorption of the samples. The schematic diagram of the laser set up is shown in Fig. 3. Two WS2-based SAs are used separately to realize passive mode locking of the fiber laser. The fiber resonator consists of a 3-m EDF (Nufen: EDFC-980-HP) with 3 dB/m absorption at 980 nm, a polarization-independent isolator, a wavelength-division multiplexer, a WS2 SA, a polarization controller, and a fiber-based optical coupler with output ratio of 5%. The other fibers and pigtails of the components are SMFs with a total length of 20.2 m (19.9 m) for SA1 (SA2). The fiber laser is pumped by a 980 nm laser diode with the maximum power of 600 mW. The dispersion parameters $D$ at 1.55 μm for EDF and SMF are $2^{16}$ ps/(nm·km) and 17 ps/(nm·km), respectively. The net cavity dispersion $\beta_2$ is calculated as about $\sim 0.38$ ps², and thus traditional soliton tends to be formed in the fiber laser.

**Experimental observations.** Continuous wave emission is achieved at an incident pump power of 10 mW using D-shaped fiber coated with WS2 (SA1). At appropriate pump power and polarization state, self-starting mode locking of fiber laser is achieved and shows clear hysteresis phenomenon. For example, mode locking starts by increasing the pump power to 25 mW while vanishes by decreasing the pump power to 13 mW. A typical mode-locking state at 300 mW is shown in Fig. 4. The central wavelength and 3-dB bandwidth of the output spectrum are measured to be 1557 nm and 2.3 nm, respectively, as shown in Fig. 4(a). Several pairs of sidebands are symmetrically distributed at both sides of the spectrum, which are the typical characteristics of standard soliton pulses.

The auto-correlation trace of the output soliton is shown in Fig. 4(b), which has a full width at half maximum of about 2.1 ps. By using a sech² fitting, the pulse duration is given as 1.32 ps from the
110 mW, the energy per pulse is calculated as 0.248 nJ. The WS2-
Considering the repetition of 8.86 MHz and average power of 600 mW, there are 50 pulses co-propagating in the fiber laser.

The signal-to-noise ratio is higher than 50 dB, indicating its good
mode-locking stability. Attributing to the peak power clamping
effect of standard soliton\(^1\), multiple pulses are formed in the fiber resonator, as demonstrated in Fig. 4(d). Noteworthily, stable single-pulse mode locking also can be achieved by decreasing the pump power to 15 mW, as shown in Fig. 4(e). The interval between adjacent pulses is measured as 113 ns, which is equal to the cavity round-trip time.

An inherent feature of the proposed SA is the ultra-high optical damage threshold. During the experiment, mode locking can be always observed when the pump power ranges from the self-starting threshold value to the maximum available pump power. Figure 5 shows the intracavity power as a function of pump power at the mode-locking state. One can observe that the intracavity power almost increases linearly with the pump. By using a linear fitting, the slope efficiency is given as 0.184. At the maximum pump power of 600 mW, mode locking operation can be maintained for several hours as long as the experiment condition keeps unchanged. Figures 7(a) and 7(b) show the spectrum and autocorrelation trace of the pulse at the pump power of 450 mW, respectively. Here, the 3-dB spectral width and pulse duration are given as 3.1 and 1.2 ps, respectively. The corresponding time bandwidth product is calculated as 0.34, suggesting that the output soliton is slightly chirped. The fundamental repetition rate is given as 8.96 MHz, as shown in the inset of Fig. 7(a). The oscilloscope trace in the inset of Fig. 7(b) indicates that multiple pulses are formed in the fiber resonator, which is similar with that of Fig. 4(d).

It is worth to note that mode locking operation can be easily established when the D-shaped fiber is coated with WS\(_2\) while vanishes when the D-shaped fiber is coated with pure-PVA, air, water, alcohol, or the mixture of water with alcohol. Additionally, we have purposely removed the WS\(_2\)-PVA film from the laser cavity to verify whether the soliton formation is purely caused by WS\(_2\). In this case, no mode locking is observed, despite that the pump power is changed from zero to the maximum (600 mW) and polarization controller is tuned over a full range. By inserting the WS\(_2\)-PVA film into fiber laser, mode locking operation can be obtained again. As a result, we conclude that the mode locking operation is purely induced by WS\(_2\) rather than other components. Based on experimental observations, we believe that WS\(_2\) nanosheets exhibit nonlinear saturable absorption property and are able to serve as mode locker for ultrafast fiber lasers.

**Discussion**

The mechanism of saturable absorption in semiconductor can be described by a two-level model\(^1\). When semiconductor is excited by light with photon energy larger than the gap energy, electrons will be transferred from the valence band to the conduction band. Under strong excitation, electrons from the valence band are excited into the conduction band and the states in the valence band become depleted, while the finial states in the conduction band are partially occupied. Further excitation from the valence band is blocked and no absorption is induced, leading to a saturable absorption condition that low-intensity (high-intensity) light experiences large (small) loss. As a result, it is easy to understand that topological insulator and MoS\(_2\) exhibit nonlinear saturable absorption property for light with photon energy larger than the gap energy.

The direct bandgap of monolayer MoS\(_2\) is about 1.8 eV (0.688 µm) and the indirect gap of the bulk material is in the range of 0.86–
1.29 eV (1.44–0.96 μm)\(^2\). It seems that MoS\(_2\) nanosheets are beyond the application as a saturable absorption device for laser wavelength larger than 1.44 μm\(^4\). However, experimental results have revealed that MoS\(_2\) exhibits saturable absorption property at 0.4, 0.8, 1.06, 1.42, 1.55, and 2.1 μm\(^22–27\). Various theories have been proposed to interpret the intriguing experimental observations, such as defect-induced bandgap decreasing\(^{14,15}\) as well as coexistence of semiconducting and metallic states\(^{28}\). Like MoS\(_2\), our experiment demonstrates that WS\(_2\) exhibits saturable absorption property at 1.55 μm, in which the energy per photon (0.8 eV) is lower than the direct bandgap of monolayer WS\(_2\) (2 eV)\(^2\) or indirect gap of the bulk WS\(_2\) (1.34 eV)\(^2\). Until now the nonlinear saturable absorption mechanism of WS\(_2\) nanosheets has not been clearly raveled at such wavelength. As MoS\(_2\) and WS\(_2\) possess similar lattice structures and photonic properties\(^15,17,21\), one could infer that saturable absorption mechanism of WS\(_2\) may be similar with that of the MoS\(_2\) reported previously\(^22,24,25\).

**Methods**

**Measurement method.** An optical spectrum analyzer (Yokogawa AQ6370), a commercial autocorrelator (APC Pulse Check), a RF analyzer, and a digital oscilloscope are employed to characterize the laser output simultaneously.

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**Author contributions**
D.M. designed the work, performed the experiments, and wrote the paper. Y.W. and S.H. fabricated and tested the D-shaped fiber. C.M. measured the property of WS₂ nanosheets. L.H. performed the part of experiment. B.J. and X.G. carried out the data analysis and revised the manuscript. W.Z., T.M. and J.Z. contributed to the scientific discussion and considerably improved the manuscript presentation. All authors discussed the results and substantially contributed to the manuscript.

**Additional information**
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