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1. Cladding waveguides are fabricated in Yb,Na:CaF₂ by femtosecond laser inscription.
2. Low propagation losses and single-mode waveguides are obtained.
3. Modification mechanism is revealed by investigating the combed μ -Raman properties.
4. Visible cooperative up-conversion emissions are achieved in the waveguides.

Cooperative Up-converted Luminescence in Yb,Na:CaF₂ Cladding Waveguides by Femtosecond Laser Inscription

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Abstract: Cladding waveguides are fabricated in Yb,Na:CaF₂ crystal by applying femtosecond laser inscription. Waveguide properties are investigated in terms of guiding behaviors and confocal micro-Raman characterizations. In addition, under 946 nm excitation, visible cooperative up-conversion emissions at 478 nm induced by Yb³⁺ ion pairs are observed while other visible bands are detected owing to the impurities of Er³⁺ and Tm³⁺ ions.

Keywords: Femtosecond laser inscription, Cladding waveguides, Cooperative up-conversion, Yb,Na:CaF₂ crystal.

1. Introduction

Femtosecond laser inscription (FLI) has been proved to be an effective technology for optical waveguide fabrication in numerous optical materials [1-4]. Waveguide structure with a tubular cladding morphology, a central unexposed waveguide core surrounded by laser induced low-index tracks, have attracted increasing attention mainly because such structures can be tailored according to demand and the properties of the host materials can be preserved well in the guiding region [5-7]. Compact lab-on-chip devices based on cladding waveguide structures have been achieved including beam couplers or splitters, novel waveguide lasers and frequency converters [8-13].

CaF₂ crystal shows unique advantages when compared with other fluoride materials. It has wide transmission band ranging from deep ultraviolet to mid-infrared;

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it also has low refractive index and nonlinear coefficient, which can reduce the nonlinear effect of high-intensity laser pumping; It exhibits lower phonon energy that can improve the quantum efficiency of fluorescence and result in relatively low lasing threshold. It also possesses a high laser damage threshold [12-18]. All these properties make CaF₂ crystal an important doping substrate for various optical applications. The Yb³⁺ ion shows a very simple electronic level structure with the ground state (²F_{7/2}) and an excited state (²F_{5/2}), leading to puny shielded 4f electrons, which makes Yb³⁺ ions easy to interact with the lattice and neighbor ions. In consequence, so called cooperative up-conversion in visible blue regions, produced by two nearby-located Yb³⁺ ions, can be observed in Yb-doped materials [19-31]. Therefore, Yb:CaF₂ crystal is considered to be an excellent candidate for visible luminescence generation. However, it has been demonstrated that the advantages of Yb:CaF₂ are often mitigated by the formation of Yb²⁺-ions and the ion clusters. An effective solution is to introduce non-active ions (such as Na⁺) into rare-earth doped CaF₂ crystal [12,32]. Visible luminescence devices based on cladding waveguides in Yb,Na:CaF₂ crystal, which combine the compact geometric of guiding structures while maintaining the advantages of the substrate material, show promising potential for applications in information technology, color display, biomedical diagnostics and underwater optical communication [33-36]. In this work, we demonstrate the formation and up-conversion of cladding waveguides in Yb,Na:CaF₂ crystal by using FLI. The guiding performance of the waveguides are observed to be excellent. By observing the micro-Raman characteristics, the mechanism of the waveguides formation is revealed. More importantly, visible blue up-conversion of Yb³⁺ ions at 478 nm based on cooperative transition in the guiding regions are reported. Extra fluorescence bands in the violet, blue, green and red regions are also detected which are due to Er³⁺ and Tm³⁺ ion impurities.

2. Experimental Procedures

The 2.0 at% Yb³⁺-ions and 5.0 at% Na⁺-ions are incorporated into CaF₂ cubic crystal which is cut into a size of 2 mm × 10 mm × 10 mm and then optically polished. The tubular cladding structures are fabricated in the prepared Yb,Na:CaF₂ crystal by applying FLI. During the fabrication process, an ultrafast Yb-doped fiber master-oscillator power amplifier laser (IMRA FCPA μ-Jewel D400) is used as laser source, delivering 360 fs pulses with a repetition rate of 500 KHz and a center

1 wavelength of 1047 nm. The laser beam with circular polarization is focused by a
 2 microscope objective (NA=0.4) into the substrate beneath one of the $10 \times 10 \text{ mm}^2$
 3 surfaces. The sample is translated through the focused laser at a speed of 20 mm/s.
 4 The inscription power of laser beam is varied from 100 mW to 160 mW with a step of
 5 20 mW, corresponding to pulse energies varied from 200 nJ to 320 nJ with a step of 40
 6 nJ. Under these conditions, arrays of parallel tracks are inscribed below the top
 7 surface following the designed geometries so as to form claddings waveguides
 8 with diameters ranging from 20 μm to 35 μm . The central depths of these cladding
 9 structures are positioned around 100 μm below the sample surface. In order to
 10 enhance the refractive index (RI) contrast of waveguiding core compared with
 11 laser-induced tracks, the scanning process is repeated with different numbers of
 12 overlapping scans (3 scans or 10 scans) for each track. Consequently, 32 cladding
 13 waveguides (numbered as WG1-WG32) are fabricated.

14 By using a linearly-polarized diode laser at 633 nm and end-face coupling,
 15 experiments are carried out to illustrate the guiding characteristics of the waveguides.
 16 A half-wave plate is employed to control the polarizations of the incident laser. Modal
 17 profiles of these structures are detected. The RI contrast Δn is calculated roughly by
 18 the formula [37]:

$$19 \Delta n = \frac{\sin^2 \theta_m}{2n_s} \quad (1)$$

20 in which θ_m is the maximum permitted incident angle. Propagation losses are
 21 estimated by detecting the incident and output power while taking Fresnel reflection
 22 of the end facets and the coupling efficiency into account. The propagation loss α can
 23 be estimated with the following equation [38]:

$$24 P_{\text{out}} = P_{\text{in}} \cdot (1 - R)^2 \cdot e^{-\alpha L} \cdot T \quad (2)$$

25 where R is the Fresnel reflection coefficient, which is calculated to be 0.0318. L
 26 stands for the length of the waveguide. T is related to the mismatch between the pump
 27 beam mode and waveguide mode which, for single-mode waveguides, can be
 28 expressed as [38]:

$$29 T = \left(\frac{2\omega_1\omega_2}{\omega_1^2 + \omega_2^2} \right)^2 \quad (3)$$

30 where ω_1 and ω_2 are the mode width of waveguide and pump beam, respectively.
 31 Presenting single-mode guidance for all waveguides, the values of T are estimated to
 32 be 0.934, 0.988, 0.995 and 0.917 for the waveguides with diameters of 35 μm , 30 μm ,
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25 μm and 20 μm , respectively. It should be pointed out that, for multi-mode waveguides, the values of parameters related to the mode mismatch are larger than the calculated results of T due to the mode transition. Thus, the propagation losses of multi-mode waveguides should be smaller than the calculated values.

The confocal Raman properties of the cladding waveguide WG1 are further investigated with a fiber-coupled confocal microscope (alpha300 J R w iTec GmbH). The excitation laser is a diode-pumped solid-state laser (532 nm, Coherent Laser). The continuous wave laser is focused via a 50 \times objective ($NA = 0.55$). The lateral and axial resolutions of the confocal system are 500 nm and 1 μm , respectively. The Raman scattered light are dispersed by a 600 mm focal length spectrometer with 1800 grooves/mm grating (UHTS 600). The signals are eventually detected using a charge-coupled-device (CCD) thermoelectrically cooled to -60°C . In order to obtain the in-depth variation of the Raman spectra, the excitation spot is scanned continuously over the cross section of WG1 with the emission line at 321 cm^{-1} . Two dimensional (2D) mappings including the emitted intensity, peak position of the emission line and emission bandwidth are obtained. Meanwhile, for easy visualization and comparison, 1D profiles of micro-Raman are also detected.

Additionally, based on the end-face coupling system, the experiment for up-conversion emission is actualized by using a 946 nm diode-pumped solid-state laser as excitation laser. The linearly polarized beam is focused and coupled into the waveguide in combination with a dielectric mirror which has high transmittance at around 946 nm in order to increase the pump power incident into the waveguide. A half-wave plate is used as to investigate fluorescence properties in both of the two polarizations. After being separated from the residual pump with a 900 nm short pass filter, the up-conversion emissions from the waveguides and, for comparison, from the bulk are detected.

3. Results and Discussion

The fabricating parameters of 32 WGs are shown in Table 1, where the experimentally obtained mode profile patterns at wavelength of 633 nm are also depicted. These waveguides are capable of supporting both TE and TM polarizations, and the mode distributions did not exhibit significant difference, which underlines the advantage of polarization independence of the cladding structures owing to their symmetric morphologies. Strong optical confinements are obtained from all

1 waveguides and the mode distributions of WG1-WG4 have been reported elsewhere
 2 [39]. Single-mode **guidance is** obtained from structures with a diameter of 20 μm ,
 3 revealing that mode numbers are reduced along with the reduction of the waveguide
 4 diameters. Furthermore, some waveguides (WG14, WG15, WG27, WG29 and WG31),
 5 although **possessing** larger diameter, are also single-mode which is related to the
 6 smaller RI contrasts induced by relatively low inscription powers **with** compare to
 7 those produced with high laser powers. Furthermore, the single-mode performance of
 8 cladding waveguide WG27 indicates lower RI contrast of waveguides fabricated with
 9 3 scans than those fabricated with 10 scans. Such a result is also numerically proved
 10 by calculating the values of RI contrast of these waveguides (see below).
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12 Figure 1(a) shows the cross-sections of **WG1 and WG17** where it can be clearly
 13 seen that the laser-induced **damage only occurs** at the modified areas, forming distinct
 14 waveguide boundaries deeply embedded inside the sample, while the core regions and
 15 the bulk outside the claddings are without any obvious damages. The propagation
 16 losses of cladding waveguides (10 scans) obtained under **TM and TE** polarization are
 17 plotted in Fig. 1(b). The minimum value is estimated to be around 0.5 dB/cm for
 18 WG1. It can be seen clearly that, at fixed inscription power, reduced propagation
 19 losses are obtained when the guiding cores are enlarged. Meanwhile, for waveguides
 20 with **the same** diameter, the propagation losses decrease monotonously when the
 21 irradiated-laser power increasing from 100 mW to 160 mW. **Fig. 1(c) shows the RI**
 22 **contrast of waveguides with 10 scans. It can be obtained that, the waveguides**
 23 **fabricated with 100 mW laser power possess minimum RI contrast around 1.0×10^{-3}**
 24 **and the waveguides inscribed with 120 mW, 140 mW and 160 mW have similar RI**
 25 **contrast (from 1.3×10^{-3} to 1.4×10^{-3}). For waveguides with 3 scans, very similar**
 26 **variation trends of propagation losses and RI contrasts are observed. Meanwhile,**
 27 **higher guiding losses related to lower RI contrast of waveguides with 3 scans are**
 28 **obtained when compared with corresponding waveguides of 10 scans. The lowest**
 29 **guiding loss is realized in WG17 with a value of approximately 0.6 dB/cm and the**
 30 **maximum RI variation of around 1.2×10^{-3} is obtained. It can be clearly observed that**
 31 **negligible differences are observed for TE and TM polarizations.**
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33 Figure 2(a) illustrates the micro-Raman emission lines obtained from the
 34 waveguide area (point A as indicated in Fig. 2(b)) and a laser-induced track (point B)
 35 of WG1. As can be seen, the micro-Raman intensity inside the track suffers from a
 36 strong quenching. In order to get complete knowledge on the spatial distributions of
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1 the changes in the micro-Raman spectra, and hence on the corresponding
 2 microstructural changes over the whole waveguide cross-section, the spatial
 3 distribution of the intensity, spectral shift and bandwidth of the emission line are
 4 measured in 2D (as shown in Figs. 2(b)-2(d)) and 1D (Figs. 2(e)-2(g)) forms. 1D
 5 profiles are measured along the green lines crossing the waveguide indicated in Figs.
 6 2(b)-2(d). As can be seen from these figures, obvious reductions in Raman intensity,
 7 blue shifts and broadening of the emission line are spatially located at the modified
 8 volumes. These phenomena, in general, can be attributed to the creation of lattice
 9 defects and damages in these regions, which are responsible for the RI reduction in
 10 the cladding areas. Furthermore, Figs. 2(b)-2(g) also demonstrate that in the active
 11 volume of the waveguide similar Raman intensity, peak position and band width are
 12 obtained in respect to bulk, which, in general, means that the lattice structures in the
 13 guiding areas are well preserved during the FLI procedure so that good optical
 14 properties of substrate material can be expected in the waveguide cores.

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Figure 3(a) depicts the up-conversion emission spectra collected from the cladding waveguides (WG1-WG17) and the bulk, which are realized under 946 nm at room temperature with fixed excitation power. As can be seen from Fig. 3(a), the up-conversion performance improved obviously when the inscription power is increased and the guiding core is enlarged due to the reduction of propagation losses. The best performance is observed in WG1. For 3 scans, the best up-conversion emission is realized in WG17 as shown in Fig. 3(a), and the intensity of the emission is much lower than that obtained in WG1. Meanwhile, in comparison with the bulk, the emission intensities are strengthened in the waveguides, which reveals the strong optical confinement of the fluorescence emission in the guiding volumes, making these waveguides promising for integrated fluorescence devices. Further evidence can be found from the photographs of visible up-conversion emissions observed in WG1 and bulk area, as exhibited in Figs. 3(b) and (c), from which a clear intensity quenching in the bulk is observed.

To investigate the details of the guided up-conversion emission, the spectra of the fluorescence generated from WG1 is measured, as described in Fig. 4. As it can be seen, an overall increase in the intensity of all the emission lines is observed when the excitation power is increased. The emission spectra show broad band covering blue-violet, blue, green and red regions, which, in a first order approximation, can be attributed to the impurities of Er^{3+} , Tm^{3+} or other rare earths in Yb^{3+} doped substrate

since it is hard to separate the Ln^{3+} ions due to their similar chemical properties. Such phenomena have been previously demonstrated in materials such as Yb:YAG waveguide [26] and Yb:Lu₃Sc₂Ga₃O₁₂ nano-garnets [22]. The peaks observed around 408 nm and 545 nm are associated with the Er^{3+} ions transitions corresponding to $(^2\text{H}_{9/2}, ^4\text{S}_{3/2}) \rightarrow ^4\text{I}_{15/2}$, respectively. The transitions $^4\text{F}_{9/2} \rightarrow ^4\text{I}_{15/2}$ of Er^{3+} ions and $^1\text{G}_4 \rightarrow ^3\text{F}_4$ of Tm^{3+} ions cause the red band with center of 657 nm [40-43]. It is found that, although the concentrations of Er^{3+} , Tm^{3+} are low, the intensities of emission lines around 657 nm are quite strong. This is mainly because the up-conversion luminescence of the impurity ions are induced by energy transfer from the Yb^{3+} ions to Er^{3+} and Tm^{3+} , which have been proved to possess very high efficiency [44]. Emission line centered at 478 nm is induced by up-conversion process of Yb^{3+} ions pairs and Tm^{3+} ($^1\text{G}_4 \rightarrow ^3\text{H}_6$). However, the generation of this peak is dominated by cooperative up-conversion emission, which can be confirmed by several evidences. Firstly, cooperative emission has been reported previously in Yb^{3+} doped materials [22-31]. Such a phenomenon can be explained in terms of the radiative relaxation of the simultaneously excited Yb^{3+} ion pairs accompanied by the emission of a visible photon with the sum of energies, which can be expressed as $^2\text{F}_{5/2} + ^2\text{F}_{5/2} \rightarrow 2^2\text{F}_{7/2} + h\nu$. [45,46] Secondly, the cooperative emission shows fairly wide band which has never been observed in up-conversion luminescence of other rare earth ions [20,47]. Finally, it has been proved that, for the up-conversion of Tm^{3+} , the emission lines around 800 nm (corresponding to the transition from $^3\text{H}_4$ to $^3\text{H}_6$) have much stronger intensity than that induced by transition of $^1\text{G}_4 \rightarrow ^3\text{H}_6$. However, in our work, the peak around 800 nm is relatively low, which in turn, proves that the emission intensity of Tm^{3+} from $^1\text{G}_4$ to $^3\text{H}_6$ is very weak [48,49]. Consequently, it can be concluded that the energy band corresponding to 478 nm is mainly attributed to cooperative up-conversion of Yb^{3+} pairs.

4. Conclusion

Cladding waveguides in Yb,Na:CaF₂ crystal are fabricated by FLI with various parameters. With optimized inscription conditions, the fabricated structures show good guiding performance in terms of low propagation losses, single-mode guidance and polarization independence. The micro-Raman characterizations reveal that laser-induced lattice defects only occur on tracks while properties of the bulk are well

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preserved in guiding volumes. Under 946 nm excitation, the visible cooperative up-conversion emissions of Yb³⁺ ion pairs at 478 nm are achieved in the waveguides while energy transfers from Yb³⁺ ions to Er³⁺ and Tm³⁺ impurities are responsible for the other bands of emission spectra. These cladding waveguides show good potential for integrated optical circuits and miniature visible fluorescence waveguide devices.

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Figure and Table captions

Figure 1. (a) The end-face microscope images of Yb,Na:CaF₂ cladding waveguides WG1 and WG17. (b) The propagation losses and (c) the refractive index contrast of waveguides with 10 scans obtained under TM and TE polarization.

Figure 2. (a) Raman spectra obtained from the guiding area and a damage track of WG1. The spatial distributions and 1D profiles of Raman intensity ((b) and (e)), Raman shift ((c) and (f)) and bandwidth ((d) and (g)) obtained from the cross-section of WG1. 1D profiles are detected along the green lines indicated in (b), (e) and (d).

Figure 3. (a) Up-conversion emissions of waveguides (WG1-WG17) and the bulk. The photographs of the visible fluorescence generated in WG1 (b) and the bulk (c).

Figure 4. Up-conversion spectra obtained from WG1 under the 946 nm excitation.

Table 1. Mode profiles observed from the fabricated 32 waveguides (WG1-WG32);

MM and SM represent multi-mode and single-mode, respectively.

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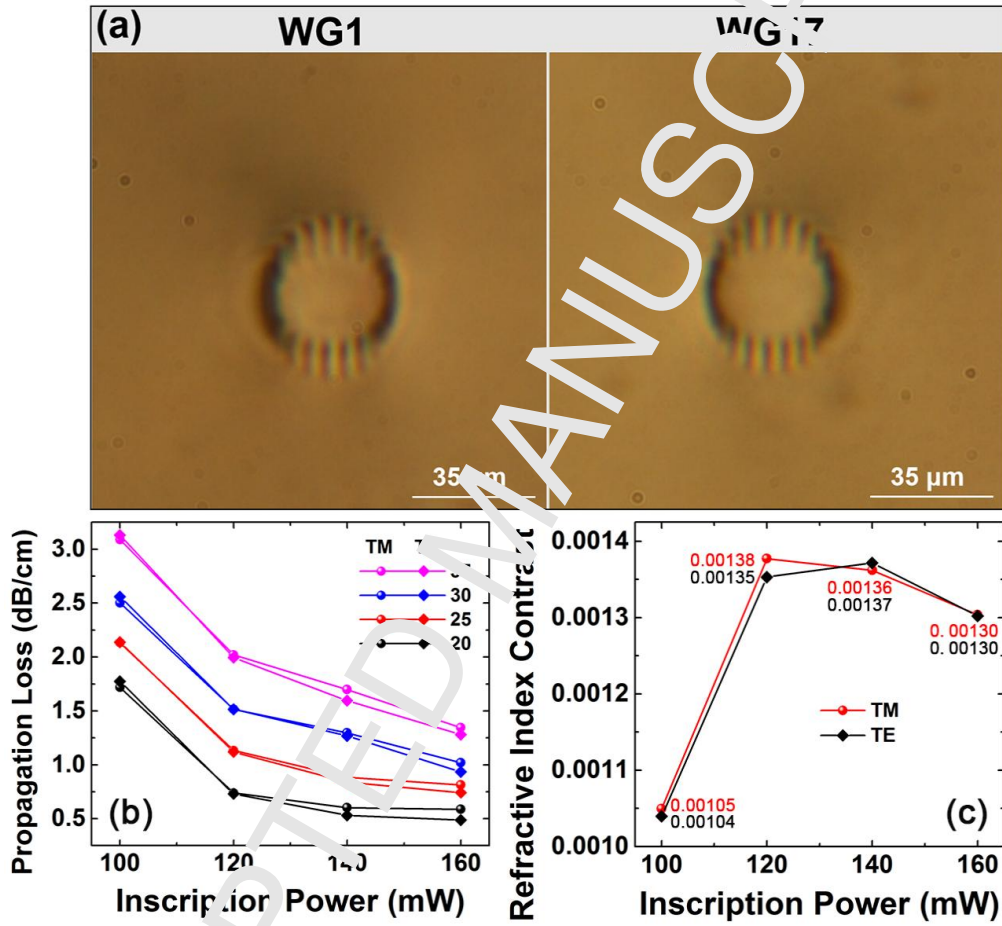


Fig. 1.

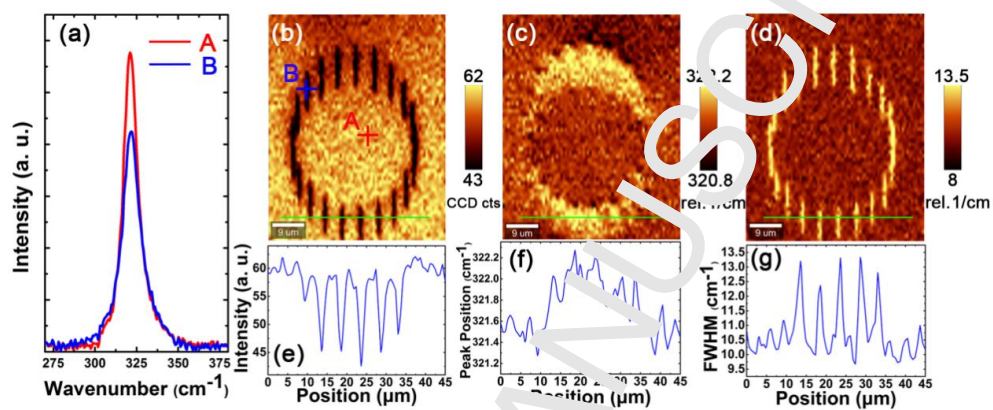


Fig. 2.

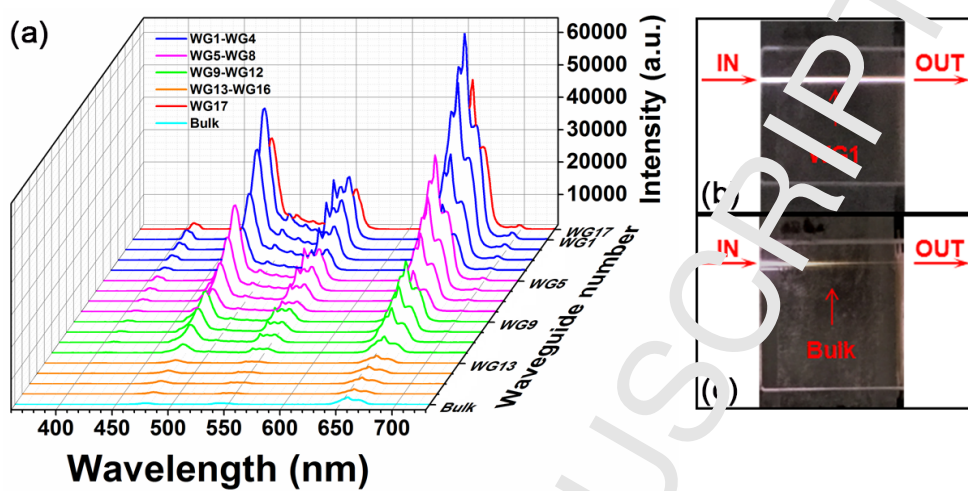


Fig. 3.

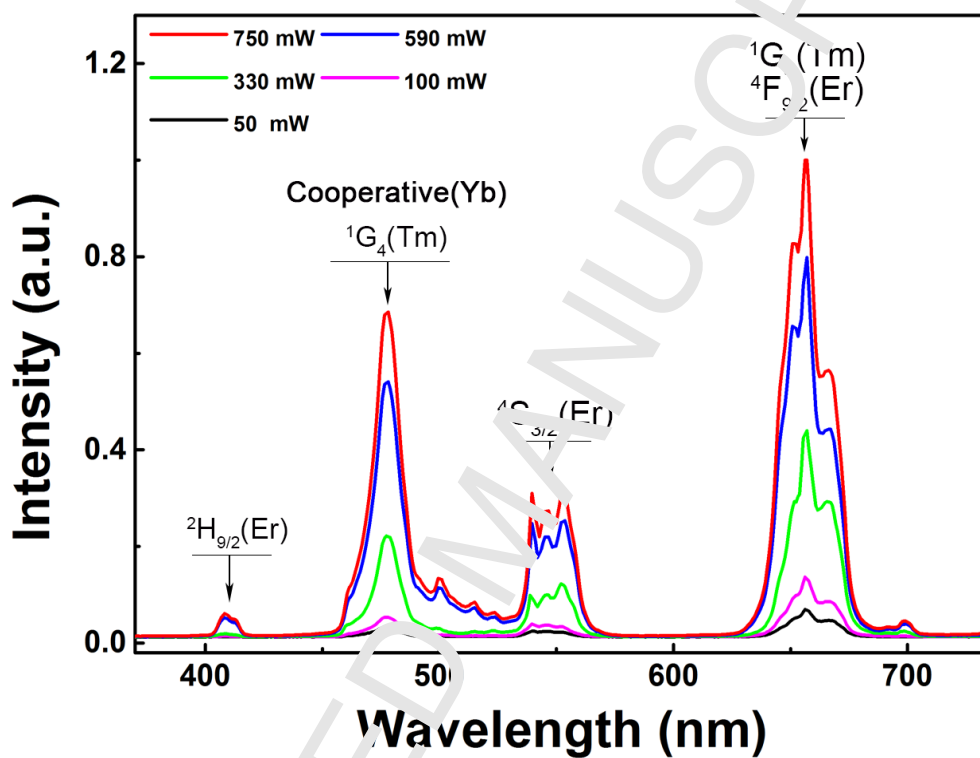


Fig. 4.

Diameter (μm)	Inscription Power (mW)	10 scans				3 scans			
		160	140	120	100	160	140	120	100
35		MM (WG1)	MM (WG5)	MM (WG9)	MM (WG13)	MM (WG17)	MM (WG21)	MM (WG25)	MM (WG29)
30		MM (WG2)	MM (WG6)	MM (WG10)	SM (WG14)	MM (WG18)	MM (WG22)	MM (WG26)	SM (WG30)
25		MM (WG3)	MM (WG7)	MM (WG11)	SM (WG15)	MM (WG19)	MM (WG23)	SM (WG27)	SM (WG31)
20		SM (WG4)	SM (WG8)	SM (WG12)	SM (WG16)	SM (WG20)	SM (WG24)	SM (WG28)	SM (WG32)

Table. 1.