Solid Optical Fiber with Tunable Bandgaps based on Curable Polymer Infiltrated Photonic Crystal Fiber

Bing Sun, Wei Wei, Chao Wang, Changrui Liao, Jing Xu, Hongdan Wan, Lin Zhang, Zuxing Zhang and Yiping Wang, Fellow, IEEE

Abstract—We demonstrated the realization and characterization of a solid photonic bandgap fiber (SPBF) with a compact size of about 10 mm and a high wavelength sensitivity of up to -4.034 nm/°C by means of fully infiltrating an ultraviolet (UV) curable polymer with a high refractive index of 1.515 into air holes of a photonic crystal fiber (PCF). To the best of our knowledge, it was the first time that the SPBF with tunable bandgaps was fabricated in the conventional index-guiding PCF. Compared with conventional fluid filled PBFS, the proposed SPBF can be stable to temperature and other environmental effects and maintain a large extinction ratio of more than 30 dB within a broad wavelength. The splicing between the SPBF and single mode fibers has been solved. Moreover, it’s observed that the bandwidth of bandgap (G2) gradually broadens with the temperature rise.

Index Terms—Microstructured optical fibers, photonic bandgaps fiber, tunable bandgaps, sensor

I. INTRODUCTION

Photonic crystal fiber (PCF) [1], a type of microstructured optical fiber (MOF) where a regular-hexagonal lattice of hollow channels is arrayed symmetrically around a central silica core, has resulted in a number of novel devices and sensing applications. In recent years, as a relatively new member of the family of MOFs, all-solid photonic bandgap fibers have attracted great attention because of their easiness to fabricate, splicing with single mode fibers, all-solid structure and promising potential application [2-6]. Note that the locations of these formed bandgaps are spectrally determined by the resonance properties of the high-index inclusions. And the high-index rods including into silica background are usually Ge-doped ones. We have to admit that the Ge-doped rods have a graded index profile with a maximum Δn of 2.03%, which forms a low index contrast, and then results in a low-extinction-ratio bandgap. Although the all-solid photonic bandgap fiber with a high index contrast (e.g. Δn=0.23) has been fabricated by the well-known stack-and-draw technique [2-3, 7], it is limited by the required pairs of glasses with closely similar softening temperatures and thermal expansion coefficients.

Fortunately, bandgaps with a high extinction ratio can be realized with infiltration of fluids into air holes of PCFs [8-9]. And the infiltration of various materials into the cladding, such as refractive index matched liquids [9], liquid crystals [8, 10-12], metals [13], ferrofluids [14-16], polymers [17-18], glasses [19-20], makes it possible to create highly tunable fiber devices. However, those fibers assisted with fluid filling are being unstable to temperature and other environmental effects. In special, the liquid surface of the liquid infiltrated PCF device at the liquid-air interface fluctuates randomly, results in a poor temporal stability.

In this paper we report the realization and characterization of a solid optical fiber by the use of a polymer material infiltrating into PCF. Note that the periodic alternating layers of dielectric material give rise to two-dimensional Bragg scattering forming a photonic band gap in the cladding. And frequencies which lie within the photonic band gap are not allowed to propagate within the cladding. The bandgaps of the cladding spontaneously depend on the refractive index of the inclusions in the PCF, i.e. the thermally tunable polymer. In fact, the SPBF can be easily fabricated owing to the polymer utilized is an ultraviolet (UV) curable liquid. Firstly, the splicing problem of the SPBF is solved according to the optimized splicing parameters. Secondly, we have numerically simulated and experimentally observed the guiding mechanism transformation between modified total internal reflection and photonic bandgap guidance. Finally, we investigate the response of the SPBF to temperature ranging from 25 °C to 95 °C.

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II. NUMERICAL SIMULATION OF BANDGAP LOCATION AND EXPERIMENTAL RESULTS

As shown in Fig. 1, a large-mode-area PCF (LAM-10, http://www.nktphotonics.com) is employed. The PCF is composed of hexagonally arrayed cladding holes with a diameter of 3.3 µm while the core is formed by omitting an air hole in the center. And the pitch of the PCF is 7.4 µm.

Fig. 1. (Left) Cross-sectional SEM images of the PCF employed; (Right) Schematic of PCF fluid filling set-up assisted with Longer Pump.

Here, we take advantage of a commercially available Norland Optical Adhesive, i.e., NOA65. The NOA65 is a clear, colorless, liquid photopolymer, whose viscosity is about 300 cps at room temperature. Frankly speaking, infiltrating NOA65 into air holes of PCFs is a complicated and time-consuming process, owing to not only its viscosity but the time required for capillary filling grows as the square of the length [21]. To solve it, we submerge one end-facet of the PCF into a liquid bottle full of NOA65, while leaving the other end-facet connected to an injection syringe to apply pressure to the fluid. As a result, the filling time can be significantly reduced owing to the existence of the pressure difference between the PCF end-facets. We utilize a commercial machine (LSP02-2A, http://www.longerpump.com.cn), where an electrically controlled moving stage has been exploited. Considering that bubbles may undesirably form and result in a discontinuous strand of fluid, the flow rate cannot be too rapid. As a result, it takes one hour for filling material to fill a ~50 mm long PCF in this experiment.

Fig. 2. Schematic of the NOA65-filled PCF. The insets side image shows the fusion joint of the NOA65-filled PCF with a single mode fiber.

Next, we focus on solving the splicing difficulty between single mode fibers (SMFs) and the fully NOA65-filled PCF. We know that the high temperature involved during fusion splicing will induce the polymer, i.e., NOA65 to boil and evaporate. As a result, it eventually leads to a bad physical strength. In the previous report [8], we take advantage of optimized splicing parameters of a commercial fusion splicer (Fujikura-60S) to implement the splicing between liquid crystal filled PCFs and SMFs. To solve it, optimized splicing parameters for the Fujikura-60S are listed in Table I. For our fiber device, the insertion losses mainly derived from splicing joints losses at the input/output and absorption loss of the polymer in the holes. In fact, the couplings between the filled and unfilled sections of the PCF can also lead to loss. Furthermore, the mode field mismatch resulting from an index guiding mode and PBG guiding mode at the splices would further bring losses.

Table I Splicing parameters for a commercial fusion splicer (Fujikura-60S)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Values</th>
<th>Units</th>
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</thead>
<tbody>
<tr>
<td>Prefusion power</td>
<td>standard-25</td>
<td>bit</td>
</tr>
<tr>
<td>Prefusion time</td>
<td>180</td>
<td>ms</td>
</tr>
<tr>
<td>Overlap</td>
<td>6</td>
<td>µm</td>
</tr>
<tr>
<td>Fusion power</td>
<td>standard-30</td>
<td>bit</td>
</tr>
<tr>
<td>Fusion time</td>
<td>300</td>
<td>ms</td>
</tr>
<tr>
<td>offset</td>
<td>-30</td>
<td>µm</td>
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Fortunately, the NOA65-filled PCF can be cured by UV light (a wavelength from 320 nm to 380 nm) with an intensity of approximately 270 mW/cm² in this experiment. In this UV-curing process, the NOA65 with a refractive index of about 1.515 is gradually transformed into a solid polymer with a refractive index of 1.524 [22]. Consequently, a solid photonic bandgap fiber (SPBF) with a compact size of about 10 mm has been fabricated. Such a compact size can probably be attributed to the higher refractive index (n=1.524) of the polymer employed.

Fig. 3. (a) First Brillouin zone and k-path for a hexagonal lattice. (b) Index profile for the single unit cell.

A full-vectorial plane wave method (FV-PWM) is applied to calculate the location of photonic bandgaps of the cladding unit cell, i.e., effective-index curves, for the fundamental mode guided in the SPBF at a temperature of 25°C. We first solve the frequency eigenvalues of the vector Helmholtz equation, and then calculate the band structure of the SPBF by considering the irreducible first Brillouin zone of the hexagonal fiber lattice, where the number of plane waves used to resolve the unit cell is 256×256 and βΛ is scanned in steps of 0.05. The first Brillouin zone of triangular lattice in cladding of SPBF is shown in Fig. 3(a), where the axis is considered to be parallel to the Γ-M direction owing to that the x-y plane is periodic. Furthermore, a single primitive unit cell, as shown in Fig. 3(b), is chosen.
Fig. 4. Calculated bandgap maps (gray color) for the SPBF and the transmission spectra of the filled (purple line) and unfilled PCF (green line). Insets: Calculated mode field distributions of the SPBF at 1290 and 1400 nm wavelength.

As shown in Fig. 4, it’s evident that three bandgaps (denoted in gray color) are observed in the calculated modal maps at the temperature of 25°C within the wavelength range from 900 to 1700 nm, where the thermo-optic effect of the pure silica background is taken into consideration in the calculations. Furthermore, the transmission spectrum of the SPBF can be measured by connecting a white-light source (NKT SuperK Compact) and an optical spectrum analyzer (YOKOGAWA AQ6370C). We also found that three bandgaps occurred within the whole wavelength range in accordance with the simulation. Some slight discrepancy in the bandgaps location is attributed to the fact that the material property of the polymer, i.e., NOA65 is not thoroughly researched. Intuitively, the operation of the SPBF may be explained in terms of antiresonant reflecting optical waveguides (ARROW), where individual rods (NOA65-filled holes) form isolated waveguides. Furthermore, the ARROW model predicts the spectral stopbands to be positioned at wavelengths corresponding to modal cutoff wavelengths of a single NOA65-filled hole. The insets shown in Fig. 4 illustrate that light at wavelengths near the cutoff wavelengths, i.e., λ=1290 nm of the individual rods’ modes can couple to these rods while the cladding becomes transparent to the light and light cannot remain confined in the core. Between cutoff wavelengths, i.e., λ=1400 nm, however, light cannot couple to the resonances of the high-index inclusions and thus remains in the core.

Fig. 5. Measured transmission spectra and the corresponding bandgap maps of the SPBF at a temperature of (a) 25°C and (b) 90°C.

We then investigate the temperature response of the SPBF by placing it in a column oven, where the temperature is raised from 25°C to 90°C in step of 10°C. As shown in Fig. 5, three bandgaps, i.e., G2, G3 and G4, appears at the temperature of 25°C, while another bandgaps, i.e., G1 and G5, located at the shortest and longest wavelength are gradually observed at the temperature of 90°C. As the temperature is increased, the edge wavelengths of the bandgap shift toward a shorter wavelength, i.e., ‘blue’ shift, due to a negative thermo-optic coefficient (-1.83×10^-4/°C) of the polymer, i.e., NOA65 [23]. As shown in Fig. 6, we tend to attribute the decline in wavelength of the G2 edges at higher temperatures to the negative thermo-optic coefficient of the NOA65 owing to that the cutoff wavelengths of the high-index inclusion’s modes shift toward a shorter wavelength with the increased temperature. Moreover, the simulation results show, in general, good qualitative agreement with the experimental measurements.

Fig. 6. The transmission spectra of the SPBF corresponding to variations in temperature ranging from 25°C to 90°C.

In order to evaluate quantitatively the temperature-induced shift of this device, we illustrate the wavelength, corresponding to -20 dB, of the left and right edges of the bandgap. This is because it is very difficult to measure the center wavelength of each bandgap. In this experiment, we characterize the bandgap 2 (G2) at different temperature for convenience’s sake. It can be seen from Fig. 7 that the left edges of G2 linearly shifted toward a shorter wavelength, i.e., ‘blue’ shift with a high sensitivity of -4.034 nm/°C while the right edges of G2 also present a ‘blue’ shift with a lower sensitivity of -1.319 nm/°C during the temperature rise. The temperature sensitivity of the reported AS-PCF is close to 2 times higher than that of the liquid-filled PCF [24]. Note that the proposed AS-PCF takes advantage over the previous results [25-26] in regard to the temperature sensitivity. As a result, the bandwidth of G2 gradual broadens with the temperature rise owing to the different sensitivity between the left and right edges of G2.

Fig. 7. Wavelengths, corresponding to a transmission of -20 dB, at the left and right edges of the G2, and the bandwidth of G2 versus the temperature.
III. CONCLUSION

In this paper, we presented a solid photonic bandgap fiber fabricated from the conventional index-guiding PCF with a curable polymer-filled cladding. The polymer inclusions introduce a high index contrast between the core and the cladding and consequently lead to lots of bandgaps. We simulated and measured the transmission spectrum from 900-1700 nm and recorded the response of the SPBF to temperature. The results illustrated our proposed SPBF is wavelength-tunable with a sensitivity of up to ~4.034 nm/C and has a large extinction ratio of more than 30 dB (G2). In conclusion, the proposed SPBF provides a new and simple highway for guidance of light and appears to extremely temporal stability.

REFERENCES


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