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HAL Id: hal-00589258
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Submitted on 28 Apr 2011

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Novel Dopants for Silica-Based Fiber Amplifiers

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Abstract: We review on some studies of the potential of new dopants in silica fibers for ultrabroadband amplification coverage in the 800-1700 nm range, including transition metal and rare-earth ions, and sensitized nano-particles.

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OCIS codes: (060.2320) Fiber optics amplifiers and oscillators; (060.2290) Fiber materials; (060.2410) Fibers, erbium

1. Introduction

Optical telecommunications systems already benefit from well developed and efficient optical fiber amplifiers operating on broad bands. Reliable, low energy consumption, broadband, flat gain erbium-doped amplifiers (EDFA) are commercially available and installed in the C-band (1535-1565 nm) and in the L-band (1565-1625 nm), where silica losses are the lowest (<0.2 dB/km). Even in the S-band (1460-1530 nm), thulium-doped fluoride glass fiber amplifiers (T DFA) are available. Although the total available amplified bandwidth is not fully used, long term basic research is of great importance in the aim of identifying innovative, cheap, reliable fiber amplifiers on the already-covered bands, and offer amplification over a broader spectral width, again using cost effective and very reliable fiber materials.

Thanks to recent improvements of silica fibers fabrication processes, their transmission window is now very wide, allowing telecommunications almost anywhere within the 800-1700 nm window, depending on the transmission distance and application. Some current telecommunication bands not yet covered by EDFA (ie unexploited or using passive systems) may in the future require amplification due to the ever growing number of users and applications. Developing active (i.e. amplified) links is therefore interesting in several bands, such as the 1st historical telecommunication window (850 nm, fiber attenuation ~ -2 dB/km), the O-band or 2nd telecommunication window (1260-1360 nm, ~ -0.5 dB/km). Even the U-band (1625-1675 nm, ~ -0.5 to -1.0 dB/km) would be exploitable (Figure 1). Alternative technologies under development are Raman fiber amplifiers (RFAs) and semiconductor amplifiers (SOAs), with some limitations including complex pumping schemes and inter-band crosstalk for RFAs, limited bandwidth per individual SOA, etc…

Besides silica-based EDFA, other glasses doped with appropriate rare-earth dopants have been proposed to cover the S and C+L bands, sometimes with good characteristics (high quantum efficiency (QE), gain flatness, gain red-shifting,…). Some are even commercially available, such as fluoride- or tellurite- or even mixed compounds oxides-glass based EDFA and TDFAs. However, many issues are still actual in fabricating cheap and reliable fibers using these materials, and fusion splicing to silica components is most of the cases impossible. Therefore silica glass is prefered as glass for cost effectiveness and reliability. The aim of this paper is to review on-going research projects on innovative amplifying dopants implemented in silica-based fibers. Issues that are usually addressed in reports are (1) identifying potential dopants and (2) examine improved manners to incorporate them in silica fibers to obtain spectrally broadened amplification. Investigations concern either conventional bands in the aim of proposing simpler architectures than existing devices, using less components, or amplification over yet uncovered bands. These research activities benefit of contributions from several science fields and technologies: process engineering, glass science, spectroscopy and high resolution material characterization.

2. Status of the silica transparent window coverage by rare earth doped amplifiers.

We review here on amplification either experimentally demonstrated and/or commercially available in the 800-1700 nm range, starting from the long wavelength side. The C- and L-bands are fully covered by silica and non-silica EDFA. Some low-phonon energy glasses (oxide and non-oxides) EDFA offer smoother and red-shifted gain curves, allowing a slight extension of the overall gain curve [1, 2]. S-band is covered only by TDFAs [3], using fluoride or other low-phonon energy glasses [1], except for silica S-band EDFA covering 1480-1520 nm, using long wavelength amplified spontaneous emission (ASE) filtering scheme [4] or gain clamping complex architecture. The O-band around 1310 nm is partially covered by praseodymium, only in low-phonon glasses [5], however the QE is low (4-6 % in fluorides) and Pr^3+ is difficult to pump (weak absorption at ~1030 nm). At shorter wavelengths, both ytterbium and neodymium ions have gain in the 1050-1150-nm range, whereas using ASE filtering [6], signal
amplification should be possible in the 980-1050-nm and 900-950-nm ranges, respectively. At last, thulium shows a narrow 20-nm gain band around 840 nm in silica [7], although a QE close to unity would be found only in low-phonon energy glasses. Therefore, the main gaps which are not covered yet, even in laboratory experiments, are approximately: 1150-1300 nm, 1350-1460 nm, 1625-1700 nm. Regarding silica-based only fiber amplifiers, these gaps are larger: 800-900 nm, 1150-1490 and 1625-1700 nm, respectively.

3. Alternative dopants with potential broad band amplification

Alternative dopants to erbium, ytterbium and neodymium in silica suffer from multi-phonon non-radiative (MP-NR) relaxations (giving low QE) and/or gain quenching due to bottlenecks on the lowest energy level of the emission transition. Two main causes induce NR decays: (1) a high material phonon energy associated with a relatively small energy gap between the dopant emitting energy level and the level just underneath (as for Tm$^{3+}$ in both 1st telecom. window and S-band) and/or (2) a strong electron-phonon coupling to the material, caused by site symmetry distortion imposed to the dopant, as for transition metals (TM). Silica has a high phonon energy (1100 cm$^{-1}$, higher than in fluoride glasses ~550 cm$^{-1}$) and offers distorted dopant sites. Therefore it is necessary to develop some kind of local environment engineering for the dopants within the silica glass network. Some examples based exclusively on silica-based doped fibers are discussed below.

Figure 1: ITU telecommunication bands and possible spectral ranges covered by standard dopants (rare-earth ions operating on high-QE transitions; top line) and alternative dopants (TM and RE ions operating on lower-QE transitions; bottom line) in silica-based fibers.

**Thulium**: Tm$^{3+}$ ion has three emission bands at 820, 1470 and 1900 nm respectively. The first two bands both originate from the same $^3H_4$ energy level, suffering strong MP-NR decay. In the 1$^{st}$ telecom. window, 11dB of gain was measured with 300 mW of 1064-nm pumping [7] whereas Cole et al. have measured only 7 dB in the S-band [8]. Both studies concerned MCVD-produced fibres. By simply codoping with aluminium, a 4-fold QE improvement was demonstrated by us and the overall lifetime decay was increased up to 60μs. Looking closer to the decay data analysis, we have shown that Tm$^{3+}$ ions were located in two very different types of sites. One of them is characterized by a almost 100 μs-lifetime (QE=15%), ever increasing when the Al-content was increased [9]. A better control of the composition and process would increase the overall QE. In the U-band, Tm$^{3+}$ has also some potential using the strong $^6F_{5/2}$=>$^4H_6$ emission band usually exploited for 1.9 μm laser emission. However, ASE filtering and gain flattening can be applied, as it was theoretically proposed for fluorides U-band TDFAs [10,11].

**Chromium**: In glasses, TM ions have potentially ten-fold spectrally larger and stronger emission cross-sections than rare-earth ions. Most importantly, their emission bands characteristics are extremely sensitive to local environment [12]. Cr$^{4+}$ in alumino-silicate fibres has a very broad band from 1000 to 1600 nm [13], although its QE is very low, due to highly distorted site symmetry[14]. By tuning the fibre composition (particularly in gallo-silicate fibers) and post-drawing heat treating, all chromium ions were stabilized into Cr$^{3+}$ in a less distorted local environment. QE as high as 10% and lifetimes equal to 20 μs were found [15]. The expected gain curve extends from 800 to 1000 nm, but necessitates visible pump wavelength, around 600-700 nm. It was recently found that the Cr$^{3+}$ ions were located in nanocrystallites grown during the heat treatment.[16].

**Bismuth**: Recently, Dvoyrin et al. obtained preliminary results of amplification (0.2 dB/m) at 1300 nm in a Bi-low-doped fiber pumped at 1064 nm at low power. The potential gain band exceeds 1150-1350 nm, as assumed from laser experiments, showing very promising 20-24 % slope efficiencies [17,18]. The exact nature of the active center responsible for this gain curve is still under study [19].
Activated nanoparticles in silica-based fibers: In order to improve some spectroscopic properties of selected amplifying ions, several research groups are trying to insert them into silica fiber via nanoparticles (NPs) selected for the local composition and site structure they would provide to the dopant. Studies performed in bulk glasses or planar waveguides are encouraging. In silica-based fibers, dielectric particles doped with Co²⁺ ions were grown by post-drawing heat-treatment in ZnO-Al₂O₃-SiO₂ core glass. Improvement of the absorption and luminescence properties were observed in the 900-nm range, but on a much narrower bandwidth than that of Cr³⁺ [20]. As mentioned above, Cr³⁺ ions were stabilized in nanoparticles in gallium-codoped silica fibers, providing luminescence in the 700-1000 nm range, more efficient and broader than in as-drawn Cr³⁺ & Cr⁴⁺ codoped fibers [16].

4. Conclusion

We have reviewed recent reports on alternative dopants for possible amplification range broadening. Although some of the studied alternative dopants suffer MP-NR decays, systematic local-environment improvement must be performed. Also new formulation and local structuration around more standard dopants are of great interest, particularly isolating ions in nanoparticles within the silica glass. Considering all possible dopants, even those of low QE may contribute to fill some spectral gaps in specific applications.

References